This Page Is Inserted by IFW Operations and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problem Mailbox.

PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶:

A61K 38/19, 39/395, 38/21, G01N 33/53

// (A61K 39/395, 38:21) (A61K 38/21,
38:19)

(11) International Patent Classification ⁶:

(43) Internation ⁶:

(44) International Patent Classification ⁶:

(45) Internation ⁶:

(46) Internation ⁶:

(47) Internation ⁶:

(48) Internation

(11) International Publication Number:

WO 96/22788

(43) International Publicati n Date:

1 August 1996 (01.08.96)

(21) International Application Number:

PCT/US96/01386

(22) International Filing Date:

26 January 1996 (26.01.96)

(30) Priority Data:

08/378,968

26 January 1995 (26.01.95) US

us

(60) Parent Application or Grant

(63) Related by Continuation

US Filed on 08/378,968 (CIP) 26 January 1995 (26.01.95)

(71) Applicant (for all designated States except US): BIOGEN, INC. [US/US]; 14 Cambridge Center, Cambridge, MA 02142 (US).

(72) Inventors; and

(75) Inventors/Applicants (for US only): BROWNING, Jeffrey, L. [US/US]; 32 Milton Road, Brookline, MA 02146 (US). MEIER, Werner [CH/US]; 31 Bedford Street, Burlington, MA 01803 (US). BENJAMIN, Christopher, D. [US/US]; 2 Oak Hill Lane, Beverly, MA 01915 (US).

(74) Agents: HALEY, James, F., Jr. et al.; Fish & Neave, 1251 Avenue of the Americas, New York, NY 10020-1104 (US).

(81) Designated States: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, ARIPO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AZ, BY, KG, KZ, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: LYMPHOTOXIN- α/β COMPLEXES AND ANTI-LYMPHOTOXIN-BETA RECEPTOR ANTIBODIES AS ANTI-TUMOR AGENTS

(57) Abstract

This invention relates to compositions and methods useful for activating LT- β receptor signaling, which in turn elicits potent anti-proliferative effects on tumor cells. More particularly, this invention relates to lymphotoxin heteromeric complexes formed between lymphotoxin- α and multiple subunits of lymphotoxin- β , which induce cytotoxic effects on tumor cells in the presence of lymphotoxin- β receptor activating agents. Also within the scope of this invention are antibodies directed against the lymphotoxin- β receptor which act as lymphotoxin- β receptor activating agents alone or in combination with other lymphotoxin- β receptor activating agents either in the presence or absence of lymphotoxin- α/β complexes. A screening method for selecting such antibodies is provided. This invention also relates to compositions and methods using cross-linked anti-lymphotoxin- β receptor antibodies either alone or in the presence of other lymphotoxin- β receptor activating agents to potentiate tumor cell cytotoxicity.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AM	Armenia	GB	United Kingdom	MW	Malawi
AT	Austria	GE	Georgia	MX	Mexico
AU	Australia	GN	Guinea	NE	Niger
BB	Barbados	GR	Greece	NL	Netherlands
BE	Belgium	HU	Hungary	NO	Norway
BF	Burkina Faso	TE	Ireland	NZ	New Zealand
BG	Bulgaria	ľŦ	Italy	PL.	Poland
BJ	Benin	JP	Japan	PT	Portugal
BR	Brazil	KR	Кепуа	RO	Romania
BY	Belanus	KG	Kyrgystan	RU	Russian Federation
CA	Canada	KP	Democratic People's Republic	SD	Sudan
CF	Central African Republic		of Korea	SE	Sweden
CG	Congo	KR	Republic of Korea	SG	Singapore
CH	Switzerland	KZ	Kazakhstan	SI	Slovenia
CI	Côte d'Ivoire	u	Liechtenstein	SK	Slovakia
CM	Cameroon	LK	Sri Lanka	SN	Senegal
CN	China	LR	Liberia	SZ	Swaziland
CS	Czechoslovakia	LT	Lithuania	TD	Chad
CZ	Czech Republic	LU	Luxembourg	TG	Togo
DE	Germany	LV	Latvia	TJ	Tajikistan
DK	Denmark	MC	Monaco	TT	Trinidad and Tobago
EE	Estonia	MD	Republic of Moldova	UA	Ukraine
ES	Spain	MG	Madagascar	UG	Uganda
FI	Finland	ML	Mali	US	United States of America
FR	Prance	MN	Mongolia	UZ	Uzbekistan
GA	Gabon	MR	Mauritania	VN	Vict Nam

LYMPHOTOXIN-α/β COMPLEXES AND ANTI-LYMPHOTOXIN-BETA RECEPTOR ANTIBODIES AS ANTI-TUMOR AGENTS

TECHNICAL FIELD OF THE INVENTION

This invention relates to compositions and methods useful for activating lymphotoxin-B receptor signaling, which in turn elicits potent anti-5 proliferative effects on tumor cells. More particularly, this invention relates to lymphotoxin heteromeric complexes formed between lymphotoxin- α and multiple subunits of lymphotoxin-B, which induce cytotoxic effects on tumor cells in the presence of lymphotoxin-ß receptor activating agents. Also within the scope of this 10 invention are antibodies directed against the lymphotoxin-B receptor which act as lymphotoxin-B receptor activating agents alone or in combination with other lymphotoxin-B receptor activating agents either in the presence or absence of lymphotoxin- α/β complexes. A 15 screening method for selecting such antibodies is provided. This invention also relates to compositions and methods using cross-linked anti-lymphotoxin-B receptor antibodies in the presence of other 20 lymphotoxin-B receptor activating agents to potentiate tumor cell cytotoxicity.

20

25

30

35

BACKGROUND OF THE INVENTION

The tumor necrosis factor (TNF) receptor family has several members whose signaling can induce tumor cell death by necrosis or apoptosis (programmed cell death).

5 The ligands TNF and lymphotoxin-α (LT-α; formerly called TNF-β) bind to and activate TNF receptors (p60 and p80; herein called "TNF-R"). TNF-R signaling initiates general immune responses to infection or stress in normal cells, but is cytotoxic to cells with transformed

10 phenotypes or to tumor cells. TNF-R signaling can selectively lyse tumor cells and virus-infected cells. The cytotoxic effects of TNF-R signaling on tumor cells are enhanced by interferon-γ (IFN-γ) and by a variety of conventional chemotherapeutic agents.

It would be useful to take advantage of the anti-proliferative or cytotoxic activities induced by TNF-R signaling in tumor cells for therapeutic purposes. However, TNF-R activation has pleiotropic effects on a variety of immunoregulatory responses including the initiation of proinflammatory cascades. Thus it has not been possible to direct the cytotoxic effects of TNF-R signaling to tumor cells without co-stimulating inflammatory responses which lead to general toxicity in humans.

Similarly, stimulation of another TNF-related receptor called the Fas receptor (FasR) can trigger cytotoxicity by programmed cell death in a variety of both tumor and non-tumor cell types. However, FasR activation has been shown to cause rapid liver necrosis, thus precluding its therapeutic application in humans.

Recently, another receptor in the TNF family called the LT- β receptor (LT- β -R) was identified (Crowe et al., Science, 264, pp. 707-10 (1994)). The LT- β -R binds heteromeric lymphotoxin complexes (LT- α/β) which comprise LT- α subunits in association with another TNF-

related polypeptide called lymphotoxin-ß (LT-ß). These LT- α /ß complexes are membrane-associated and most have a LT- α 1/ß2 stoichiometry (Browning et al., <u>Cell</u>, 72, pp. 847-56 (1993); Browning et al., <u>J. Immunol.</u>, 154, pp. 33-46 (1995)).

5

25

30

35

By analogy to TNF-R and other TNF-like receptors, the activation of LT-B-R signaling is thought to occur when multiple receptors on the cell surface are brought into close proximity (Crowe et al., Science, 264, 10 pp. 707-10 (1994)). This process is referred to as receptor clustering. The TNF and LT ligands are multivalent complexes which can simultaneously bind to and thus aggregate more than one receptor. Receptor clustering as a means for receptor activation in other 15 systems has been well-documented, especially for receptor tyrosine kinases (Ullrich and Schlessinger, Cell, 61, pp. 203-212 (1990); Kolanus et al., Cell, 74, pp. 171-83 (1993)). Accordingly, administering LT-α1/β2 ligands and/or LT-B-R activating agents which can induce the 20 clustering and downstream signaling of LT-B-R molecules on the surface of target tumor cells would be useful for directly stimulating the LT-B-R pathway in these cells.

Signaling by LT-ß-R, like TNF-R, can activate pathways that lead to cytotoxicity and cell death in tumor cells. Importantly, LT- α 1/ β 2 ligands do not bind to TNF-R with any significant affinity. For this reason, directed LT- β -R activation in tumor cells would trigger cytotoxicity in those cells without stimulating the inflammatory pathways associated with TNF-R activation. Treatment with LT- α 1/ β 2 and/or other LT- β -R activating agents would thus be useful for treating or reducing the advancement, severity or effects of tumorigenic cells (neoplasia) while overcoming the potent side effect problems which have been encountered when TNF-R or FasR activation has been tried as an anti-tumor treatment.

- 4 -

SUMMARY OF THE INVENTION

The present invention solves the problems referred to above by providing pharmaceutical compositions and methods for treating tumor cells by stimulating LT-B-R signaling without co-stimulating TNF-5 R-associated inflammatory responses. In one embodiment, lymphotoxin complexes formed between LT-α and multiple subunits of LT-β are provided (LT-α/β heteromeric complexes) which induce cytotoxic effects on cells bearing the LT-B-R in the presence of a LT-B-R activating 10 agent. The preferred compositions and methods of this embodiment comprise LT- α 1/ β 2 complexes in the presence of a LT- β -R activating agent. More preferably, the LT- α 1/ β 2 complexes are in a soluble rather than a membrane-bound form, and the LT-B-R activating agent is IFN-v. 15

In another embodiment of the invention, at least one antibody directed against LT-\$\beta\$-R (anti-LT-\$\beta\$-R Ab) is used as a second LT-\$\beta\$-R activating agent in conjunction with the LT-\$\alpha\$/\$\beta\$ heteromeric complex. The preferred compositions and methods of this embodiment are characterized by LT-\$\alpha\$/\$\beta\$2 in the presence of IFN-\$\gamma\$ as a first activating agent, and at least one anti-LT-\$\beta\$-R Ab as a second LT-\$\beta\$-R activating agent. More preferably, the LT-\$\alpha\$/\$\beta\$2 complexes are soluble and the antibody is a monoclonal antibody (anti-LT-\$\beta\$-R mAb).

20

25

30

In another embodiment of the invention, at least one anti-LT- β -R Ab in the presence or absence of a second LT- β -R activating agent is used without an exogenous LT- α/β heteromeric complex. The preferred compositions and methods of this embodiment comprise at least two anti-LT- β -R monoclonal antibodies (anti-LT- β -R mAbs) which recognize non-overlapping epitopes of LT- β -R in combination with IFN- γ .

In a further embodiment, this invention
provides pharmaceutical compositions and methods for

- 5 -

potentiating tumor cell cytotoxicity characterized by cross-linked anti-LT-\(\theta\)-R Abs used in conjunction with a second LT-\(\theta\)-R activating agent. In one preferred embodiment, individual anti-LT-\(\theta\)-R Abs are immobilized by cross-linking them onto a surface. In another preferred embodiment, the anti-LT-\(\theta\)-R Abs are cross-linked in solution. More preferably, the anti-LT-\(\theta\)-R Abs are monoclonal antibodies and the second LT-\(\theta\)-R activating agent is IFN-\(\gamma\).

This invention further provides a novel screening process for selecting LT-β-R activating agents, such as anti-LT-β-R Abs, that function in combination with LT-α/β heteromeric complexes to promote tumor cell death. The assay makes use of the increased sensitivity of human adenocarcinoma cells to LT-α/β heteromeric complexes in the presence of LT-β-R activating agents in a cytotoxicity assay. The procedure used to test putative LT-β-R activating agents is exemplified for the case of anti-LT-β-R antibodies, and comprises the following steps:

- 1) Tumor cells (e.g., HT29 human adenocarcinoma cells) are cultured for several days in media containing IFN- γ and purified LT- α 1/ β 2 in the presence or absence of the particular anti-LT- β -R Ab being assayed;
- 2) The cells are treated with a dye that stains living cells; and

25

3) The number of stained cells is quantitated to determine the fraction of tumor cells killed in the presence of the LT-α1/β2, IFN-γ and the test anti-LT-β-R Ab in each sample. Alternatively, the number of surviving cells can be determined by any of a number of well-known assays which measure cell viability, such as ³H-thymidine incorporation into DNA.

An anti-LT- β -R Ab (or an Ab combination) which significantly increases the percentage of tumor cells killed in this assay is a LT- β -R activating agent within the scope of this invention. This cytolytic assay can be adapted to identify new LT- β -R activating agents which function in combination with LT- α/β heteromeric complexes.

BRIEF DESCRIPTION OF THE DRAWINGS

- Figure 1A. Sizing analyses of the purified LT-α/β
 10 heteromeric complex forms separated by TNF-R and LT-β-R immunoaffinity chromatography. Purified proteins were analyzed on a TSK 3000 HPLC resin in phosphate-buffered saline. The position of various size markers is shown.
- Figure 1B. Ion exchange analyses of purified LT forms using a Poros carboxymethyl column (4.6mm x 100 mm) on a BioCad instrument (Perceptive Biosystems). 27 μg of each sample protein was loaded onto a column and eluted in a gradient of 0 to 1M NaCl over 20 column volumes at 5 ml/min in a Buffer containing 16.66 μM Hepes, 16.66 μM Na Acetate, and 16.66 μM Mes buffer (pH 6.5).
- Figure 2A. Comparison of the cytotoxic activity of:
 Anti-Fas receptor mAb CH-11 (-Φ-); TNF (-O-); LT-α (-□-);
 LT-α1/β2 (-■-); and LT-α2/β1 (-Φ-) on human
 25 adenocarcinoma HT29 cells in either the presence or
 absence of 80 U/ml IFN-γ.
- Figure 2B. Comparison of the ability of 5 μg/ml of human IgG (-Φ-), soluble p60TNF-R-Fc (-O-) and soluble LT-β-R-Fc receptor-immunoglobulin chimeras (-D-) to inhibit the cytotoxic effects in Fig. 2A in the presence of 80 U/ml IFN-y.

- 7 -

Figure 3. Anti-LT-β-R mAbs potentiate the cytotoxic effect of LT-α1/β2 on human adenocarcinoma HT29 cells.

(A) The LT-α1/β2 cytolytic effects on HT29 cells are potentiated by the presence of anti-LT-β-R mAb CDH10.

5 LT-α1/β2 effects were measured without mAb (-•-), and in the presence of 0.5 μg/ml control IgG1 (---), 0.05 μg/ml CDH10 (-0-) and 0.5 μg/ml (---) CDH10. (B) The LT-α1/β2 cytolytic effects on HT29 cells are inhibited by the presence of the anti-LT-β-R mAb BDA8. LT-α1/β2 effects

10 were measured in the presence of 2 μg/ml control IgG1 (---) or anti-LT-β-R mAb BDA8 (---). The difference between the behavior of the CDH10 and BDA8 anti-LT-β-R mAbs in this assay is one indication that they are directed to different epitopes of the LT-β-R.

- Figure 4. Immobilized anti-LT-ß-R mAbs are cytotoxic to human adenocarcinoma HT29 cells. (A) The anti-LT-ß-R mAbs have a direct cytotoxic effect on HT29 cells when they are immobilized on a surface. Plates were coated with IgG1 (-•-), a mAb directed against an unrelated abundant cell surface antigen HT29/26 (-•-), BDA8 (-•-) and CDH10 (-□-). (B) The effects of soluble anti-LT-ß-R mAbs alone on the growth of HT29 cells. Symbols as in (A). These anti-LT-ß-R mAbs in their soluble form do not have significant cytotoxic effects on HT29 cells when administered individually.
 - Figure 5. Representative quantitation of the enhanced cytotoxicity to tumor cells by treating with pairs of soluble anti-LT-B-R mAbs. (A) The cytotoxic effects on HT29 cells of control IgG1 (100 ng/ml), anti-LT-B-R mAb BHA10 (100 ng/ml), anti-LT-B-R mAb CBE11 (50 ng/ml), BHA10 (100 ng/ml) + IgG (100 ng/ml), and BHA10 (100 ng/ml) + CBE11 (50 ng/ml). IFN-y was present at 80U/ml.

 (B) The cytotoxic effects on HT29 cells of control IgG1

30

(100 ng/ml), anti-LT-B-R mAb CDH10 (100 ng/ml), anti-LT-B-R mAb CBE11 (50 ng/ml), CDH10 (100 ng/ml) + IgG1 (100 ng/ml), and CDH10 (100 ng/ml) + CBE11 (50 ng/ml).

IFN-Y was present at 80U/ml. (C) The cytotoxic effects on HT29 cells of control IgG1 (100 ng/ml), anti-LT-B-R mAb CDH10 (33 ng/ml), anti-LT-B-R mAb AGH1 (50 ng/ml), and CDH10 (33 ng/ml) + AGH1 (50 ng/ml) on HT29 cells IFN-Y was present at 80U/ml. (D) As in (C), except that WiDr human adenocarcinoma cells were used in the cytolytic assay (Raitano and Korc, J. Biol. Chem., 265, pp. 10466-472 (1990)).

Figure 6. Tumor size in SCID mice treated with an anti-LT-B-R mAb. (A) Size of the human adenocarcinoma WiDr tumor in SCID mice 30 days after inoculation with an antibody co-treatment. Mice were treated on days 1 and 2 15 with saline, IFN-y alone, an anti-LT-B-R mAb (CBE11) with and without IFN-y and a control anti-human LFA-3 mAb (1E6) with IFN-y. The mean of each group is indicated by a crossbar. Means, standard deviations, and number of 20 animals (in parentheticals) for the five groups (left to right) were: 0.88 +/- 0.59 (14), 1.21 +/- 0.7 (21), 0.041+/- 0.052 (16), 0.11 +/- 0.1 (12), and 0.98 +/- 1.16 (12). (B) Size of the human adenocarcinoma WiDr tumor in SCID mice from 14 to 49 days after tumor cell 25 inoculation with a 15 day post-inoculation antibody treatment. Tumors were grown to an average diameter of 0.53 cm (0.076 cc) without any treatment and i.p. injections were started on day 15 and continued as indicated by the arrows. Means and standard deviations 30 are indicated for a group of 12 animals treated either with IFN- γ alone (1 x 10° U/injection) (- \Box -), IFN- γ with 50 µg 1E6 anti-LFA-3 mAb (-0-), IFN-y with 50 µg CBE11 anti-LT-&-R mAb (-A-) or 50 µg CBEll anti-LT-&-R mAb alone (not shown).

DETAILED DESCRIPTION OF THE INVENTION

5

10

25

30

In order that the invention herein described may be fully understood, the following detailed description is set forth.

The term "anti-tumor activity" refers to the ability of a substance or composition to block the proliferation of, or to induce the death of tumor cells which interact with that substance or composition.

The term "apoptosis" refers to a process of programmed cell death.

The term "cytotoxic activity" refers to the ability of a substance or composition to induce the death of cells which interact with that substance or composition.

15 The term "epitope" (or antigenic determinant) is defined as the part of a molecule that combines with a single antigen binding site on an antibody molecule. A single epitope is recognized by a monoclonal antibody (mAb). Multiple epitopes are normally recognized by polyclonal antibodies (Ab).

The "Fc domain" of an antibody refers to a part of the molecule comprising the CH2, CH3 and hinge regions but lacking the antigen binding sites.

The term "interferon inducing agent" refers to any agent which is capable of directly or indirectly stimulating the endogenous production of either type I (IFN- α , IFN- β) or type II (IFN- γ) interferons. Examples of interferon inducing agents include double stranded RNA molecules, and a variety of plant or pharmaceutically-derived compounds.

The terms "LT- α mutein" and "LT- β mutein" refer to LT- α or LT- β polypeptides having one or more amino acid changes compared to the amino acid sequence of the corresponding native polypeptide.

- 10 -

The term "LT- β -R activating agent" refers to any agent which can augment ligand binding to LT- β -R, cell surface LT- β -R clustering or LT- β -R signaling, or which can influence how the LT- β -R signal is interpreted within the cell. Examples of LT- β -R activating agents include IFN- α , IFN- γ , TNF, interferon inducing agents, soluble anti-LT- β -R Abs, cross-linked anti-LT- β -R Abs and multivalent anti-LT- β -R Abs.

The term "LT-ß-R signaling" refers to all molecular reactions associated with the LT-ß-R pathway and subsequent molecular reactions which result therefrom.

15

30

The term "anti-LT-\$-receptor antibody" ("anti-LT-\$-R Ab") refers to any antibody that recognizes and binds to at least one epitope of the LT-\$ receptor.

The term "anti-LT-B receptor monoclonal antibody" ("anti-LT-B-R mAb") refers to any monoclonal antibody that recognizes and binds to a single epitope of the LT-B-R.

The term "cross-linked anti-LT-ß-R (m)Abs"
refer to antibodies directed against the LT-ß-R which
have either been cross-linked to each other to form
antibody agglomerates in solution using an anti-LT-ß-R
antibody (Ab) or (mAb) cross-linking agent, or which have
been immobilized in close proximity to one another on a
surface or matrix.

The term "anti-LT-ß-R Ab (or mAb) cross-linking agent" refers to any agent which can covalently or non-covalently aggregate anti-LT-ß-R Abs in solution so that the Abs can bind to and potentiate target cell surface LT-ß receptor clustering. Such cross-linking agents include but are not limited to chemical cross-linking agents, secondary antibodies which react with portions of the anti-LT-ß-R Abs or mAbs, and soluble or surface-bound

Fc receptors -- either endogenous or added exogenously -- which can bind to anti-LT-B-R Abs.

The terms "LT- α biological activity", "LT- β biological activity", and "LT- α/β biological activity" are defined as: 1) immunological cross-reactivity with an antibody directed against at least one epitope of the corresponding native subunit or complex of subunits; or 2) the ability of the LT subunit or complex of subunits to compete for ligand binding sites on a LT-specific receptor such as TNF-R or LT- β -R; or 3) having the ability to stimulate an immune regulatory response or cytotoxic activity qualitatively in common with a native LT subunit or complex.

5

10

25

30

35

The term "LT-α/ß heteromeric complex" refers to

a stable association between at least one LT-α subunit
and more than one LT-ß subunits. The subunits can
associate through electrostatic, van der Waals, or
covalent interactions. Preferably, the LT-α/ß
heteromeric complex has at least two adjacent LT-ß

subunits and lacks adjacent LT-α subunits. Most
preferably, the complex has the stoichiometry LT-α1/β2.

The term "multivalent ligand" refers to a molecule or complex which has more than one receptor binding site and which is capable of simultaneously binding and bringing into close proximity at least two receptor molecules.

A "type I leader sequence" is an amino-terminal portion of a eukaryotic protein which serves as a signal to direct the protein to the endoplasmic reticular (ER) membrane and often through the entire secretion pathway. The leader sequence is usually cleaved off by a signal peptidase in the ER membrane.

A "signal sequence" is the functional equivalent of a eukaryotic type I leader sequence in prokaryotic hosts, and directs the translocation of

proteins into or across lipid bilayer membranes of a bacterium.

5

10

25

30

A "soluble LT- α/β heteromeric complex" is a LT- α/β heteromeric complex comprising soluble LT- β subunits, wherein the amino acid sequences which localize the polypeptide to the membrane have been deleted or inactivated, rendering the LT- β subunit soluble. Soluble LT- α/β heteromeric complexes can be secreted by an appropriate host cell which has been engineered to express both subunits.

A "surface LT- α/β complex" is a complex comprising LT- α and membrane-bound LT- β subunits which is displayed on the cell surface.

Production of Membrane-bound LT- α/δ Complexes

Cell surface lymphotoxin complexes have been characterized in CD4* T cell hybridoma cells (II-23.D7) that express high levels of LT (Browning et al., J. Immunol., 147, pp. 1230-37 (1991); Androlewicz et al., J. Biol. Chem., 267, pp. 2542-47 (1992)). Mature

LT-α lacks a transmembrane domain and is localized to the cell surface through interaction with at least one membrane-bound LT-ß subunit. Membrane-bound (surface) LT-α/ß heteromeric complexes have predominantly a LT-α1/β2 stoichiometry.

LT- β as a cell membrane protein binds LT- α during synthesis, thus "targeting" the LT- α to the cell membrane. In the absence of LT- β , LT- α is secreted into the extracellular medium. LT subunits normally assemble into complexes inside the cell prior to protein export into the membrane. Once LT- β subunits are inserted into the membrane, they do not form stable complexes with secreted LT- α . Thus if the membrane-bound form of a LT- α/β heteromeric complex is desired, it is preferable

to co-express the desired LT- α and LT- β subunits within the same cell.

The surface LT- α/β heteromeric complex can be reconstructed by co-transfection of host cells with both the LT- α and LT- β genes. Surface LT complexes cannot be observed on stable cell lines which express either LT gene alone. However, if the host cell normally produces large amounts of LT- α (e.g. RPMI 1788 cells; see below), then transfection with a LT- β gene which encodes the desired LT- β polypeptide should be sufficient to generate LT- α/β complexes comprising full-length LT- α subunits.

5

10

15

30

35

Co-expression of LT- α and LT- β polypeptides in a number of eukaryotic expression systems leads to their assembly and export as active ligand (Crowe et al., <u>J. Immunol. Methods</u>, 168, 79-89 (1994)). Host systems that can be used include but are not limited to CHO cells, COS cells, B cells including myelomas, baculovirus-infected insect cells and yeast.

The LT-α subunit of the LT-α/ß heteromeric

complexes of this invention can be selected from
lymphotoxin-α, native human or animal lymphotoxin-α,
recombinant lymphotoxin-α, soluble lymphotoxin-α,
secreted lymphotoxin-α, lymphotoxin-α muteins having LT-α
biological activity, or lymphotoxin-α fragments of any of
the above having LT-α biological activity.

The LT- α polypeptide can be any soluble form of the molecule including active fragments thereof which can be produced in eukaryotic expression systems, wherein the natural LT- α leader sequence will be cleaved off. Alternatively, fusions of the mature LT- α sequence with a heterologous signal sequence can be used to maximize the secretion of LT- α in other host systems. Signals are chosen based on the intended host cell, and may include bacterial, yeast, mammalian and viral sequences. The native signal, or the vascular cell adhesion molecule-1

- 14 -

(VCAM-1) signal sequence is suitable for use in mammalian expression systems.

LT- α polypeptides can also be fused to polypeptides having a prolonged plasma half-life such as immunoglobulin chains or fragments thereof. Plasma proteins which may be used to enhance plasma half-life include serum albumin, immunoglobulins, apolipoproteins, and transferrin. Polyethylene glycol (PEG) attachment may stabilize the polypeptide and lower its immunogenicity. Preferably the LT- α fusion protein is not significantly immunogenic in the subject to be treated and the plasma protein does not cause undesirable side effects in subjects due to its normal biological activity.

5

10

15

20

25

30

Human LT- α is glycosylated on N and O residues, and depending on the source, exhibits considerable sugarbased microheterogeneity. The oligosaccharide composition of the particular LT- α chosen to form the LT complex may affect in vivo clearance rates (Fukushima et al., Arch. Biochem. Biophys., 304, pp. 144-53 (1993)). Since glycosylation variants can be produced by expression in different host cells, this is one factor to be considered in selecting a source of LT- α .

LT- α can be purified from a B lymphoblastoid line RPMI 1788, which constitutively secretes LT- α and which can be induced to secrete higher levels by treating with the phorbol ester PMA (Aggarwal et al., J. Biol. Chem., 259, pp. 686-91 (1984)). Alternatively, the cloned human LT- α gene can be used to recombinantly produce LT- α polypeptides in different host systems including bacteria (Schoenfeld et al., J. Biol. Chem., 266, pp. 3863-69 (1991)); baculovirus-infected insect cells (Crowe et al., J. Immunol. Methods, 168, pp. 70-89 (1994)); and mammalian cells (Browning and Ribolini, J.

- 15 -

Immunol., 143, pp. 1859-67 (1989); Fukushima et al., Arch. Biochem. Biophys., 304, pp. 144-53 (1993)).

Portions of the LT- α gene which encode polypeptide fragments having LT- α biological activity can be evaluated using routine screening assays. Useful screening assays for LT- α biological activity include competitive inhibition assays with native LT- α bound to TNF-R, or measuring either directly or indirectly by inhibition the ability of the LT- α to induce cytotoxicity of tumor cells in assays known to the art. Preferably, LT- α fragments are assembled into heteromeric complexes with LT- β and the complexes assayed for LT- α/β biological activity by competitive inhibition with LT- α/β bound to LT- β -R, or for their ability to induce cytotoxicity of tumor cells in the assays disclosed herein.

10

15

20

25

30

35

Lymphotoxin-ß, also referred to as p33, has been identified on the surface of T lymphocytes, T cell lines, B cell lines and lymphokine-activated killer cells. LT-ß is the subject of applicants' co-pending international applications PCT/US91/04588, published January 9, 1992 as WO 92/00329; and PCT/US93/11669, published June 23, 1994 as WO 94/13808, which are herein incorporated by reference.

The LT-ß gene encodes a polypeptide of 240-244 amino acids (Browning et al., <u>Cell</u>, 72, pp. 847-56 (1993)). LT-ß is a type II membrane protein with a short N-terminal cytoplasmic domain followed by a membrane anchoring domain of 30 hydrophobic amino acids. It has a single N-linked glycosylation site and has only one cysteine residue which does not appear to be involved in intersubunit disulfide bond formation.

The LT-ß subunits comprising the LT- α/β heteromeric complexes of the present invention can be selected from lymphotoxin- β , native human or animal lymphotoxin- β , recombinant lymphotoxin- β , soluble

lymphotoxin-B, secreted lymphotoxin-B, lymphotoxin-B muteins having LT-B biological activity, or lymphotoxin-B fragments of any of the above having LT-B biological activity.

As discussed above for the LT- α polypeptide, the LT- β polypeptides can also be modified to increase their solubility or plasma half-life using the same methods. Likewise, portions of the LT- β gene which encode polypeptide fragments having LT- β biological activity can be evaluated using routine screening assays as discussed for LT- α .

Production of soluble complexes

5

10

25

30

Soluble (non-membrane-bound) LT-\alpha/\beta heteromeric complexes comprise LT-\beta subunits which have been changed from a membrane-bound to a soluble form. These complexes are described in detail in applicants' co-pending international application (PCT/US93/11669, published January 9, 1992 as WO 94/13808). Soluble LT-\beta peptides are defined by the amino acid sequence of lymphotoxin-\beta wherein the sequence is cleaved at any point between the end of the transmembrane region (i.e. at about amino acid #44) and the first TNF homology region (i.e. at amino acid #88) according to the numbering system of Browning et al., Cell, 72, pp. 847-56 (1993).

Soluble LT-ß polypeptides may be produced by truncating the N-terminus of LT-ß to remove the cytoplasmic tail and transmembrane region (Crowe et al., Science, 264, pp. 707-710 (1994)). Alternatively, the transmembrane domain may be inactivated by deletion, or by substitution of the normally hydrophobic amino acid residues which comprise a transmembrane domain with hydrophilic ones. In either case, a substantially hydrophilic hydropathy profile is created which will reduce lipid affinity and improve aqueous solubility.

Deletion of the transmembrane domain is preferred over substitution with hydrophilic amino acid residues because it avoids introducing potentially immunogenic epitopes.

The deleted or inactivated transmembrane domain

may be replaced with or attached to a type I leader
sequence (e.g. the VCAM-1 leader) such that the protein
is secreted beginning with a sequence anywhere from
between val40 to pro88. Soluble LT-ß polypeptides may
include any number of well-known leader sequences at the
N-terminus. Such a sequence would allow the peptides to
be expressed and targeted to the secretion pathway in a
eukaryotic system. See, e.g., Ernst et al., United
States Patent No. 5,082,783 (1992).

Soluble LT-α/ß heteromeric complexes may be

produced by co-transfecting a suitable host cell with DNA encoding LT-α and soluble LT-ß (Crowe et al., J. Immunol. Methods, 168, pp. 79-89 (1994)). Soluble LT-ß secreted in the absence of LT-α is highly oligomerized. However, when co-expressed with LT-α, a 70 kDa trimeric-like structure is formed which contains both proteins. It is also possible to produce soluble LT-α1/β2 heteromeric complexes by transfecting a cell line which normally expresses only LT-α (such as the RPMI 1788 cells discussed above) with a gene encoding a soluble LT-ß polypeptide.

LT- α and LT- β polypeptides may be separately synthesized, denatured using mild detergents, mixed together and renatured by removing the detergent to form mixed LT heteromeric complexes which can be separated (see below).

Purification of LT-α1/82 Complexes

30

 $Soluble\ LT-\alpha 1/\beta 2\ heteromeric\ complexes\ are$ separated from co-expression complexes comprising a different subunit stoichiometry by chromatography using

PCT/US96/01386

TNF and LT- β receptors as affinity purification reagents. The TNF receptors only bind within α/α clefts of LT complexes. The LT- β receptor binds with high affinity to β/β clefts, and with lower affinity to α/β clefts of heteromeric LT- α/β complexes. Accordingly, LT- α 3 and LT- α 2/ β 1 will bind to TNF-R. The LT- β -R can also bind LT- α 2/ β 1 trimers (within the α/β clefts) but cannot bind LT- α 3. In addition, the LT- β -R (but not TNF-R) binds LT- α 1/ β 2 and LT- β n (the exact composition of such preparation is unknown, however, they are large aggregates).

The receptor affinity reagents can be prepared as either a soluble extracellular domain (see for example Loetscher et al., J. Biol. Chem., 266, pp. 18324-29

(1991)), or as chimeric proteins with the extracellular ligand binding domain coupled to an immunoglobulin Fc domain (Loetscher et al., J. Biol. Chem., 266, pp. 18324-29 (1991); Crowe et al., Science, 264, pp. 707-710 (1994)). Receptors are coupled to affinity matrices by chemical cross-linking using routine procedures.

There are two schemes by which the LT- α 1/ β 2 ligand can be purified using receptors and immunoaffinity chromatography. In the first scheme, a supernatant from an appropriate expression system coexpressing both LT- α and the truncated LT- β form is passed over a TNF-R column. The TNF-R will bind LT- α 3 and LT- α 2/ β 1 trimers. The flow through from the TNF-R column will contain LT- β (n) and LT- α 1/ β 2.

25

In the second scheme, all LT-β-containing forms

(LT-β(n), LT-α1/β2 and LT-α2/β1) are bound to and eluted from a LT-β-R column using classical methods such as chaotrophe or pH change. (LT-α3 flows through this column). The eluate is neutralized or the chaotrophe removed, and the eluate is then passed over a TNF-R column, which binds only to the LT-α2/β1 trimers. The

flow through of this column will contain LT-B(n) and LT- $\alpha 1/B2$ trimers.

In both cases, pure LT- α 1/ β 2 trimers can be separated from LT- β by subsequent gel filtration and/or ion exchange chromatographic procedures known to the art.

Alternatively, different forms of LT- α/β heteromeric complexes can be separated and purified by a variety of conventional chromatographic means. It may also be preferable to combine a series of conventional purification schemes with one of the immunoaffinity purification steps described above.

Source of Anti-LT-8-R Antibodies

5

10

25

30

Polyclonal antibody sera directed against the human LT-B receptor are prepared using conventional techniques by injecting animals such as goats, rabbits or mice subcutaneously with a human LT-B receptor-Fc fusion protein (Example 2) in complete Freund's adjuvant, followed by booster intraperitoneal or subcutaneous injection in complete Freunds. Polyclonal antisera containing the desired antibodies which are directed against the LT-B receptor are screened by conventional procedures.

Mouse monoclonal antibodies (mAbs) directed against a human LT-ß receptor-Fc fusion protein are prepared by intraperitoneal immunization of RBF mice repetitively with a CHO cell-derived recombinant LT-ß receptor-Fc fusion protein (LT-ß-R-Fc) attached to protein A sepharose beads in the absence of adjuvant. Animals are finally boosted with soluble LT-ß-R-Fc (both i.p. and i.v.), spleen cells are fused using classical protocols, and hybridomas are screened by ELISA (Ling et al., J. Interferon and Cytokine Res., 15, pp. 53-59 (1995)). Hybridomas are further screened for their ability to block binding of activated II-23 hybridoma

PCT/US96/01386 WO 96/22788

- 20 -

cells -- which express surface LT- α 1/ β 2 -- to LT- β -R-Fccoated plates in a cell panning assay. Pure mAbs are prepared by protein A sepharose purification of IgG from hybridoma culture supernatants.

5

20

25

30

35

Various forms of anti-LT-B-R antibodies can also be made using standard recombinant DNA techniques (Winter and Milstein, Nature, 349, pp. 293-99 (1991)). For example, "chimeric" antibodies can be constructed in which the antigen binding domain from an animal antibody is linked to a human constant domain (e.g. Cabilly et 10 al., US 4,816,567; Morrison et al., Proc. Natl. Acad. Sci. U.S.A., 81, pp. 6851-55 (1984)). Chimeric antibodies reduce the observed immunogenic responses elicited by animal antibodies when used in human clinical treatments. 15

In addition, recombinant "humanized antibodies" which recognize the LT-B-R can be synthesized. Humanized antibodies are chimeras comprising mostly human IgG sequences into which the regions responsible for specific antigen-binding have been inserted (e.g. WO 94/04679). Animals are immunized with the desired antigen, the corresponding antibodies are isolated, and the portion of the variable region sequences responsible for specific antigen binding are removed. The animal-derived antigen binding regions are then cloned into the appropriate position of human antibody genes in which the antigen binding regions have been deleted. Humanized antibodies minimize the use of heterologous (inter-species) sequences in human antibodies, and are less likely to elicit immune responses in the treated subject.

Construction of different classes of recombinant anti-LT-B-R antibodies can also be accomplished by making chimeric or humanized antibodies comprising the anti-LT-B-R variable domains and human constant domains (CH1, CH2, CH3) isolated from different classes of immunoglobulins. For example, anti-LT-B-R IgM antibodies with increased antigen binding site valencies can be recombinantly produced by cloning the antigen binding site into vectors carrying the human μ chain constant regions (Arulanandam et al., J. Exp. Med., 177, pp. 1439-50 (1993); Lane et al., Eur. J. Immunol., 22, pp. 2573-78 (1993); Traunecker et al., Nature, 339, pp. 68-70 (1989)).

In addition, standard recombinant DNA

techniques can be used to alter the binding affinities of recombinant antibodies with their antigens by altering amino acid residues in the vicinity of the antigen binding sites. The antigen binding affinity of a humanized antibody can be increased by mutagenesis based on molecular modeling (Queen et al., Proc. Natl. Acad. Sci. U.S.A., 86, pp. 10029-33 (1989); WO 94/04679).

It may be desirable to increase or to decrease the affinity of anti-LT-\$B-R Abs for the LT-\$B-R depending on the targeted tissue type or the particular treatment schedule envisioned. For example, it may be advantageous to treat a patient with constant levels of anti-LT-\$B-R Abs with reduced ability to signal through the LT-\$B pathway for semi-prophylactic treatments. Likewise, anti-LT-\$B-R Abs with increased affinity for the LT-\$B-R may be advantageous for short-term, tumor-targeted treatments.

20

25

Screening Anti-LT-8-R Antibodies For LT-8-R Activating Agents

The anti-LT- β -R antibodies of this invention can potentiate the anti-tumor activity of LT- α/β heteromeric complexes (preferably LT- α 1/ β 2) in the presence of an LT- β -R activating agent such as IFN- γ . These anti-LT- β -R antibodies are also referred to herein

as LT- β -R activating agents. The antibodies which act as LT- β -R activating agents are selected as follows:

- 1) A series of tissue culture wells containing tumor cells such as HT29 cells are cultured for three to four days in media containing a LT- β -R activating agent such as IFN- γ , and purified LT- α/β heteromeric complex -- preferably LT- $\alpha1/\beta2$ -- in the presence or absence of serial dilutions of the anti-LT- β -R Ab being tested;
- 10 2) A vital dye stain which measures
 mitochondrial function such as MTT is added to the cell
 mixture and reacted for several hours;
- 3) The optical density of the mixture in each well is quantitated at 550 nm wavelength light (OD 550).
 15 The OD 550 is inversely proportional to the number of tumor cells killed in the presence of the LT-α/β heteromeric complex, the LT-β-R activating agent and test anti-LT-β-R Ab in each well.

The preferred antibodies of this invention

which act individually as LT-β-R activating agents in the presence of LT-α1/β2 and IFN-γ include the BKA11, CDH10, BHA10 and BCG6 anti-LT-β-R mAbs (Table 2, infra).

Cross-linking Anti-LT-8-R Antibodies

The cross-linked anti-LT-β-R antibodies of this
invention act individually as LT-β-R activating agents
without exogenous LT-α/β heteromeric complexes in the
presence of a second LT-β-R activating agent such as
IFN-γ. Cross-linked anti-LT-β-R Abs apparently bind to
and induce clustering of cell surface LT-β receptors
thereby activating LT-β receptor-mediated targeted cell
death.

In one embodiment, one or more types of anti-LT-B-R antibodies are cross-link d by immobilization onto a water-insoluble matrix or surface. Derivatization with

a bifunctional agent is useful for cross-linking the antibodies to the water-insoluble support matrix or surface. Agents commonly used to effect cross-linking of antibodies to a water insoluble support matrix or surface include 1,1 bis(-diazoacetyl)-2-phenylethane, glutyraldehyde, N-hydroxysuccinamide esters including esters with 4-azidosalicylic acid, homobifunctional imidoesters including disuccinimidyl esters, and bifunctional maleimides such as bis-N-maleimido-1,8octane. Derivatizing agents such as methyl-3-[(p-10 azidophenyl) dithio] propioimidate form photoactivatable intermediates which can be selectively cross-linked when stimulated with light. Reactive water-insoluble matrices such as cyanogen bromide-activated carbohydrates and the substrates described in United States Patent Nos. 15 3,959,080; 3,969,287; 3,691,016; 4,195,128; 4,247,642; 4,229,537; 4,055,635; and 4,330,440 can also be used for protein immobilization and cross-linking.

The surfaces to which the antibodies are attached can be non-proteinaceous polymer, usually a 20 hydrophilic polymer either from natural or synthetic sources. Hydrophilic polyvinyl polymers such as polyvinylalcohol (PVA) and polyvinylpyrrolidone (PVP) can be used. Also useful are polyalkylene ethers such as polyethylene glycol, polypropylene glycol, 25 polyoxyethylene esters or methoxy polyethylene glycol, polyoxyalkylenes such as polyoxyethylene and polyoxypropylene, and block copolymers of polyoxyethylene and polyoxypropylene (Pluronics); polymethacrylates; carbomers; branched or unbranched polysaccharides which 30 comprise the saccharide monomers D-mannose, D- and Lgalactose, fucose, fructose, D-xylose, L-arabinose, Dglucuronic acid, sialic acid, D-galacturonic acid, D-mannuronic acid (e.g. poly-mannuronic acid or alginic acid), D-glucosamine, D-galactosamine, D-glucose and 35

PCT/US96/01386 WO 96/22788

- 24 -

neuraminic acid including homopolysaccharides and heteropolysaccharides such as lactose, amylopectin, starch, hydroxyethyl starch, amylose, dextran sulfate, dextran, dextrins, glycogen, or the polysaccharide subunit of acid mucopoly-saccharides, e.g. hyaluronic acid; polymers of sugar alcohols such as polysorbitol and polymannitol; and heparin or heparon.

5

20

30

The polymer prior to cross-linking is preferably water soluble and preferably contains only a singly reactive chemical group to avoid multiple cross-10 linking events with the antibody. In any case, reaction conditions should be optimized to reduce cross-linking and to recover products -- either directly or by a subsequent gel filtration or chromatographic step -having a substantially homogenous molecular weight range. 15 The optimal molecular weight of the cross-linked antibody matrix will be determined by routine experimentation using the cytotoxicity and receptor binding assays disclosed herein.

The final conjugate after cross-linking is preferably soluble in physiological fluids such as blood. The polymer should not be highly immunogenic in the conjugate form, and should possess a viscosity compatible with intravenous infusion or injection if either is an intended route of administration. 25

The polymer may also be water insoluble. Materials which may be used include hydrophilic gels, or shaped articles having surfaces to which the antibodies can be immobilized such as surgical tubing, catheters, or drainage conduits. It is preferable to use solid support materials which are biologically compatible and substantially inert in physiological surroundings. A material is biologically compatible if it does not substantially stimulate immune responses including

inflammation, or attract fibrotic cells when placed inside the body of a subject.

5

10

30

35

Anti-LT-ß-R Abs may also be immobilized onto surfaces which have been covalently or non-covalently coated with secondary antibodies that will bind to the primary anti-LT-ß-R Abs (e.g., goat anti-mouse IgG antibodies; see Example 7). Each anti-LT-ß-R mAb tested individually, when immobilized onto a surface with secondary antibodies, acts as an LT-ß-R activating agent in the presence of IFN-y (Figures 4 and 7).

In an alternative embodiment, cross-linked anti-LT-B-R Abs in solution act as LT-B-R activating agents. Anti-LT-B-R Abs can be cross-linked by means of an anti-LT-B-R Ab (or mAb) cross-linking agent. An anti-LT-B-R Ab (or mAb) cross-linking agent according to 15 this invention is any agent capable of either covalently linking, or of non-covalently aggregating the anti-LT-B-R Abs (or mAbs) in solution so that the cross-linked anti-LT-B-R Abs (or mAbs) can bind to and potentiate target cell surface LT-B-R clustering. Such anti-LT-B-R Ab (or 20 mAb) cross-linking agents include but are not limited to chemical cross-linking reagents which can be reacted with the antibodies in a controlled manner as described above. Alternatively, secondary antibodies, Sepharose A, Fc receptors, or other agents that will bind to and 25 aggregate multiple primary anti-LT-B-R Abs without blocking their activity can be used to form anti-LT-B-R Abs agglomerates in solution.

Multiple Anti-LT-8-R Abs In Solution Act As LT-8-R Activating Agents

Compositions comprising multiple anti-LT-B-R

Abs in solution which act as LT-B-R activating agents by
potentiating surface LT-B-R clustering are provided by
this invention. Polyclonal anti-LT-B-R Abs directed
against different epitopes of the LT-B-R can be used.

25

Preferably, the anti-LT-ß-R Abs are monoclonal Abs directed against different and non-overlapping epitopes of the LT-ß-R.

The combined anti-LT-B-R mAb approach to LT-B receptor activation requires combining two non-overlapping epitopes. Moreover, it is likely that productive receptor aggregation is only achieved with certain epitopes. We have identified the presence of at least four unique LT-B-R immunoreactive epitopes.

Additional epitopes (as defined by new mAbs) may be identified by continuing to fuse immunized mouse spleen cells, by immunizing different species of animals, and by using different routes of immunization.

Epitopes can also be directly mapped by
assessing the ability of different mAbs to compete with
each other for binding to the LT-B-R using BIAcore
chromatographic techniques (Pharmacia BIAtechnology
Handbook, "Epitope Mapping", Section 6.3.2, (May 1994);
see also Johne et al., J. Immunol. Methods, 160, pp.
191-8 (1993)).

Individual LT-B-R mAbs can be grouped into at least four classes according to their ability to cooperate in combination with other LT-B-R mAbs in killing tumor cells in cytolytic assays (Example 8; Table 1). For example, the BDA8 mAb in Group I does not act in combination with the AGH1 mAb in Group I to promote tumor cell cytotoxicity. Likewise, the Group III BKA11 and CDH10 mAbs do not cooperate in a tumor cell cytotoxicity assay.

Figure 5A-C shows the effects of administering representative anti-LT-β-R mAbs alone and in pairwise combination to tumor cells in cytotoxicity assays in the presence of IFN-γ as a LT-β-R activating agent. The Group IV anti-LT-β-R mAb CBE11 used alone has a slight cytotoxic effect which is enhanced in combination with

the Group II mAb BHA10 (Figure 5A). CBE11 elicits a similar effect with th Group III mAb CDH10 (Figure 5B).

The cytotoxicity caused by administering a combination of anti-LT-B-R mAbs in solution is not peculiar to the HT29 tumor cell line. Figure 5C shows that the Group I AGH1 mAb and the Group III CDH10 mAb act synergistically in killing two different tumor cell lines (HT29 cells and WiDr cells) derived from human adenocarcinoma tumors.

10 Summary of Anti-LT-8-R mAbs Characteristics

All of the anti-LT-ß-R mAbs of this invention, when cross-linked by immobilization, act as LT-ß-R activating agents in the presence of a second LT-ß-R activating agent such as IFN- γ . The ability of anti-LT-ß-R mAbs to act as LT-ß-R activating agents in the presence or absence of LT- α 1/ß2 in solution often varies according to the state of the cells at the time of the test. Table 2, infra, summarizes the properties of the anti-LT-ß-R mAbs characterized by this invention.

The Group I mAbs BDA8 and AGH1 do not function as LT-β-R activating agents in solution with LT-α1/β2. The BDA8 mAb actually blocks the anti-tumor effect of LT-α1/β2 (Figure 3B and Table 2). In contrast, the Group II anti-LT-β-R mAbs BCG6 and BHA10 have mixed agonistic and antagonistic effects when administered with LT-α1/β2. The Group III anti-LT-β-R mAbs BKA11 and CDH10 are unique in their ability to act as LT-β-R activating agents which potentiate anti-tumor effects in the presence of LT-α1/β2 and a second LT-β-R activating agent such as IFN-γ without exhibiting the antagonistic effects often seen with the Group II mAbs BCG6 and BHA10.

It is important to keep in mind that the classification of the anti-LT-B-R mAbs based on their ability to cooperate in tumor cell cytolytic assays

10

15

35

reflects that they interact with distinct epitopes of the LT- β -R. The mAbs comprising a single group do not, however, necessarily have the same binding affinities for their cognate epitopes. Thus the variable results seen when comparing the effects of different mAbs belonging to the same or different groups may represent differences in binding affinities. Accordingly, it is possible that a Group I or a Group IV mAb with a higher binding affinity for the LT- β -R could be isolated which would function like the Group III mAbs as a LT- β -R activating agent in the presence of LT- α 1/ β 2.

The hybridoma cell lines or subclones thereof which produce the anti-LT-B-R mAbs described above were deposited on January 12, 1995 with the American Type Culture Collection (ATCC) (Rockville, MD) according to the provisions of the Budapest Treaty, and were assigned the ATCC accession numbers designated as follows:

20	a) b) c) d) e) f)	CELL LINE AG.H1.5.1 BD.A8.AB9 BC.G6.AF5 BH.A10 BK.A11.AC10 CB.E11.1	mAb Name AGH1 BDA8 BCG6 BHA10 BKA11 CBE11	ATCC Accession No. HB 11796 HB 11798 HB 11794 HB 11795 HB 11799 HB 11793
25	g)	CD.H10.1	CDH10	нв 11797

All restrictions on the availability to the public of the above ATCC deposits will be irrevocably removed upon the granting of a patent on this application.

30 Anti-LT-8-R IgM Monoclonal Antibodies Function as LT-8-R Activating Agents

Anti-LT-B-R mAbs which comprise more than the usual two IgG antigen binding sites will also function in solution as cell surface LT-B-R cross-linking agents, and will accordingly fall within th definition of a LT-B-R

activating agent according to this invention. The antigen binding sites of an anti-LT-B-R mAb can be built into IgM molecules -- which have ten antigen binding sites -- using standard recombinant DNA and hybridoma techniques (Example 12).

Alternatively, one may collect and enrich for complete mouse (or other animal) IgM molecules isolated by hybridoma fusion techniques after a single immunization with antigen. One way to enrich for IgM molecules would be to immunize CD40 signaling-deficient mice (Kawabe et al., Immunity, 1, pp. 167-78 (1994); Xu et al., Immunity, 1, pp. 423-31 (1994)). These mice cannot effectively produce IgGs and therefore their response to challenge by antigen is enriched for IgM isotypes.

10

15

20

25

Anti-LT-B-R IgM antibodies, by virtue of their increased valency, can effectively aggregate LT-B-R molecules within the plane of the membrane, thereby enhancing LT-B-R signaling as compared to their IgG counterparts having two antigen binding sites. A dramatic example of the increased efficiency of multivalent antibodies in receptor clustering is seen with antibodies to the Fas receptor, where the IgM form is very potent and normal bivalent IgGs are not effective in solution (Yonihara and Yonihara, J. Exp. Med., 169, pp. 1747-56 (1989); Alderson et al., Int. Immunol., 6, pp. 1799-1806 (1994)).

Likewise, the apo-1 mAb to the Fas receptor is an IgG3 mAb. This mAb is a potent cytotoxic agent which relies on Fc interactions unique to IgG3 subtypes to aggregate into larger polyvalent forms. Removal of the Fc region creates a F(ab)₂ form that cannot associate into larger aggregates and which is inactive (Dhein et al., <u>J. Immunol.</u>, 149, pp. 3166-73 (1992)). Thus by analogy, it

is predicted that IgM versions of anti-LT-B-R mAbs will be potent anti-tumor agents.

Anti-LT-8-R mAbs Inhibit Tumor Growth in Mice

The ability of a LT-B-R activating agent such as an anti-LT-B-R mAb to inhibit human tumor cell growth 5 in vitro (Examples 6-8 and 13) may be indicative of in vivo anti-tumorigenic activity. Experiments performed in immunodeficient (SCID) mice demonstrate that an anti-LT-B-R mAb (CBE11) can efficiently block tumor formation by human adenocarcinoma WiDr cells (Example 14; Figure 10 6). Mice inoculated subcutaneously (s.c.) with WiDr cells form measurable tumors within two weeks. When mice were treated i.p. with the CBEll mAb at the same time as the WiDr cells were inoculated s.c., tumor outgrowth was dramatically blocked (Figure 6A). The anti-tumor action 15 of the CBE11 anti-LT-B-R mAb was enhanced by adding IFN-y; CBE11 was effective, however, even without exogenous IFN-y. In the CBE11 + IFN-y group, 7 of 16 animals completely lacked tumors, whereas the remaining animals had small nodules that had not progressed at 2 20 months. The CBE11 alone treated mice were similar to the CBE11 + IFN-y group at 30 days. The CBE11 alone treated mice, however, eventually developed slowly growing tumors. There were statistically significant differences between the CBE11 (+/- IFN-y) groups and the control 25 groups (saline, IFN-y alone and control anti-human LFA-3 mAb (1E6) + IFN-γ), but no significant differences among the control groups. The 1E6 and CBE11 mAbs are both IgG1 antibodies. The 1E6 mAb effectively coats the tumor cells but does not block tumor growth. Thus complement 30 or natural killer cell-mediated events are not the sole basis for the anti-tumor activity of the CBE11 anti-LT-B-R mAb .

30

35

The efficacy of the CBE11 mAb in inhibiting tumor growth in vivo in the absence of xogenous IFN-y was unexpected since there was a dependence on IFN-y for measurable LT-B-R-based in vitro cytotoxic effects.

5 Either there is some crossover of mouse IFN- γ onto human IFN- γ receptors, or other mechanisms may be involved in vivo.

The CBE11 anti-LT-B-R mAb can also inhibit growth of an established tumor in mice (Figure 6B). Mice were inoculated s.c. with WiDr human adenocarcinoma cells 10 at day 1 and tumors were allowed to develop for 15 days (Example 14). Tumors in animals treated i.p. with IFN-y alone, or with the control anti-human LFA-3 mAb (1E6) + IFN-y, continued to increase in size over the course of the 7-week experiment. In contrast, tumors treated with 15 the CBE11 anti-LT-B-R mAb (+ IFN-y or alone) stopped growing, and following three injections of CBEll antibody over a three week period, tumor growth was arrested out to 49 days post-inoculum when the experiment was 20 terminated (Figure 6B).

These experiments demonstrate that an anti-LT-B-R mAb which activates LT-B-R signaling can effectively inhibit tumor formation at early stages and can also block continued tumor cell growth at later stages of tumorigenesis in vivo. These experiments also demonstrate that administration of a single LT-B-R activating agent may be effective for treating or reducing the advancement, severity or effects of neoplasia in an affected animal.

The procedures described in Example 14 may be used to identify LT-B-R activating agents according to this invention which function alone or in combination to inhibit tumor cell growth <u>in vivo</u>. It is envisioned that other LT-B-R activating agents -- including but not limited to those identified using <u>in vitro</u> tumor cell

10

15

20

25

30

cytotoxicity assays -- may have similar anti-tumor effects in vivo when administered either alone or in combination to animals or humans.

The Use of IFN-y and Other LT-6-R Activating Agents

The cytotoxic effects of LT- α/β heteromeric complexes and of cross-linked or multiple anti-LT- β -R Abs on tumor cells is enhanced by the presence of a LT- β -R activating agent, particularly IFN- γ . Human adenocarcinoma cells of intestinal origin (HT29 cells) have previously been shown to be sensitive to FasR signaling (Yonehara and Yonehara, J. Exp. Med., 169, pp. 1747-56 (1989)); and to TNF and LT- α in the presence of IFN- γ (Browning et al., J. Immunol., 143, pp. 1859-67 (1989)).

The amount of LT- β -R activating agent required to enhance the anti-tumor activity of LT- α/β heteromeric complexes, anti-LT- β -R Abs or other LT- β -R activating agents of this invention will depend on the cell or tissue type being treated, and also with the mode of treatment, and can be determined empirically using routine procedures. The LT- β -R activating agent can be provided at a concentration or delivered at a rate determined to be effective in conjunction with other LT- β -R activating agents administered, taking into consideration the factors listed above.

Alternatively, endogenous LT-β-R activating agents such as interferons like IFN-γ, which may be produced by cells or tissue surrounding the target tumor cells, can be relied upon. Endogenous IFN-γ is normally produced upon viral infection, and is also found in the vicinity of tumors (Dinge et al., Immunity, 1, pp. 447-56 (1994)).

Any agent which is capable of inducing interferons, preferably IFN-y, and which potentiates the

15

20

25

cytotoxic effects of LT- α/β heteromeric complexes and anti-LT- β -R mAbs on tumor cells falls within the group of LT- β -R activating agents of this invention. While virus infection normally induces IFN- γ production, the levels of endogenous IFN- γ may be enhanced by other agents (Example 10). For example, clinical experiments have demonstrated interferon induction by double stranded RNA (dsRNA) treatment. Accordingly, polyriboguanylic/polyribocytidylic acid (poly-rG/rC) and other forms of dsRNA are effective as interferon inducers (Juraskova et al., Eur. J. Pharmacol., 221, pp. 107-11 (1992)).

The interferon stimulator from Glycyrrhiza glabra (Acharya et al., Indian J. Med. Res., 98, pp. 69-74 (1993)), and pharmaceutical agents, many of which are orally administrable, may also be used to boost endogenous interferon levels. Such interferon inducers include: imiquimod (Bernstein et al., Antiviral Res., 20, pp. 45-55 (1994)); saparal (Paramonova et al., Vopr. Virusol., 39, pp. 131-34 (1994)); aryl pyrimidones such as bropirimine (Onishi and Machida, Hinyokika Kiyo, 40, pp. 195-200 (1994)); Ridostin (Cheknev et al., Vopr. Virusol., 39, pp. 125-28 (1994)).

Several of these interferon inducing agents have been characterized as inducers of type I interferons such as IFN- α . Type I interferons can also function as LT- β -R activating agents but are less potent than IFN- γ .

Treatments Using LT- α/β Complexes and LT- β -R Activating Agents

The compositions of this invention will be

administered at an effective dose to treat the particular clinical condition addressed. Determination of a preferred pharmaceutical formulation and a therapeutically efficient dose regiment for a given application is well within the skill of the art taking into consideration, for example, the condition and weight

of the patient, the extent of desired treatment and the tolerance of the patient for the treatment.

Typically, humans can tolerate up to 100-200 $\mu g/m^2$ of TNF before serious toxicity is manifested (Schiller et al., <u>Cancer Res.</u>, 51, pp. 1651-58 (1991)). 5 In mice, dosages in the range of 1-5 µg/mouse/day given with 5x104 units of recombinant human IFN-y caused human primary tumor regression (Balkwill et al., CIBA Foundation Symposium (1987); Havell et al., J. Exp. Med., 167, pp. 1067-85 (1988)). Based on the relative 10 effectiveness of TNF and LT- α 1/ β 2 in the HT29 cytolytic assays, approximately 5-25 μg/mouse/day of LT-α1/β2 will provide a therapeutic dose range. Extrapolating to the human, it is expected that LT- α 1/ β 2 dosages of at least 1 mg/m^2 will be required in combination with an LT-B-R 15 activating agent such as IFN-y.

Historically, IFN-γ therapy has been undertaken either at maximum tolerated doses in the range of 100-250 μg/m², or at "immunomodulatory" levels in the range of 10-25 μg/m² (see e.g. Kopp et al., J. Immunother., 13, pp. 181-90 (1993)). Combination therapies with two interferons have used 4x106 units/m² of IFN-α and approximately 250 μg/m² of IFN-γ (Niederle et al., Leuk. Lymphoma, 9, pp. 111-19 (1993)). Intermediate doses of about 25-100 μg/m² of IFN-γ in combination with the LT-α/β heteromeric complexes or purified anti-LT-β-R-Abs described herein are expected to be suitable starting points for optimizing treatment doses.

Administration of the LT-α/β heteromeric

complexes and cross-linked anti-LT-β-R Abs of this invention, including isolated and purified forms of the antibodies or complexes, their salts or pharmaceutically acceptable derivatives thereof, may be accomplished using any of the conventionally accepted modes of

- 35 -

administration of agents which exhibit anti-tumor activity.

10

15

20

25

30

The pharmaceutical compositions used in these therapies may also be in a variety of forms. These include, for example, solid, semi-solid and liquid dosage forms such as tablets, pills, powders, liquid solutions or suspensions, suppositories, and injectable and infusible solutions. The preferred form depends on the intended mode of administration and therapeutic application. Modes of administration may include oral, parenteral, subcutaneous, intravenous, intralesional or topical administration.

The LT-α/ß heteromeric complexes, IFN-γ, and anti-LT-β-R Abs may, for example, be placed into sterile, isotonic formulations with or without cofactors which stimulate uptake or stability. The formulation is preferably liquid, or may be lyophilized powder. For example, the LT complexes and/or anti-LT-β-R Abs and IFN-γ may be diluted with a formulation buffer comprising 5.0 mg/ml citric acid monohydrate, 2.7 mg/ml trisodium citrate, 41 mg/ml mannitol, 1 mg/ml glycine and 1 mg/ml polysorbate 20. This solution can be lyophilized, stored under refrigeration and reconstituted prior to administration with sterile Water-For-Injection (USP).

The compositions also will preferably include conventional pharmaceutically acceptable carriers well known in the art (see for example Remington's Pharmaceutical Sciences, 16th Edition, 1980, Mac Publishing Company). Such pharmaceutically acceptable carriers may include other medicinal agents, carriers, genetic carriers, adjuvants, excipients, etc., such as human serum albumin or plasma preparations. The compositions are preferably in the form of a unit dose and will usually be administered one or more times a day.

The pharmaceutical compositions of this invention may also be administered using microspheres, liposomes, other microparticulate delivery systems or sustained release formulations placed in, near, or otherwise in communication with affected tissues or the 5 bloodstream. Suitable examples of sustained release carriers include semipermeable polymer matrices in the form of shaped articles such as suppositories or microcapsules. Implantable or microcapsular sustained release matrices include polylactides (U.S. Patent No. 10 3,773,319; EP 58,481), copolymers of L-glutamic acid and gamma ethyl-L-glutamate (Sidman et al., Biopolymers, 22, pp. 547-56 (1985)); poly(2-hydroxyethyl-methacrylate) or ethylene vinyl acetate (Langer et al., J. Biomed. Mater. Res., 15, pp. 167-277 (1981); Langer, Chem. Tech., 12, 15 pp. 98-105 (1982)).

Liposomes containing LT-\alpha/\beta heteromeric complexes and/or anti-LT-\beta-R Abs and IFN-\gamma can be prepared by well-known methods (See, e.g. DE 3,218,121;

20 Epstein et al., Proc. Natl. Acad. Sci. U.S.A., 82, pp. 3688-92 (1985); Hwang et al., Proc. Natl. Acad. Sci. U.S.A., 77, pp. 4030-34 (1980); U.S. Patent Nos. 4,485,045 and 4,544,545). Ordinarily the liposomes are of the small (about 200-800 Angstroms) unilamellar type in which the lipid content is greater than about 30 mol.\gamma cholesterol. The proportion of cholesterol is selected to control the optimal rate of LT complex and/or anti-LT-\beta-R Abs and IFN-\gamma release.

The LT-α/ß heteromeric complexes and anti
10 LT-β-R Abs of this invention may also be attached to liposomes containing other LT-β-R activating agents, chemotherapeutic agents or IFN-y to supplement the IFN-y typically found in the region of tumors. Attachment of LT complexes and anti-LT-β-R Abs to liposomes may be accomplished by any known cross-linking agent such as

5

25

30

heterobifunctional cross-linking agents that have been widely used to couple toxins or chemotherapeutic agents to antibodies for targeted delivery. Conjugation to liposomes can also be accomplished using the carbohydrate-directed cross-linking reagent 4-(4-maleimidophenyl) butyric acid hydrazide (MPBH) (Duzgunes et al., J. Cell. Biochem. Abst. Suppl. 16E 77 (1992)).

Advantages of therapeutic compositions based on anti-LT-B-R activation

An anti-tumor therapy based upon LT-B-R 10 activation would have several advantages. LT-B-R binds to LT- α/β heteromeric complexes with high affinity in β/β clefts, and with lower affinity in α/β clefts created at the interfaces between adjacent LT- α and LT- β subunits. 15 In contrast, the TNF receptors bind to $LT-\alpha/\beta$ heteromeric complexes with high affinity only in α/α clefts. Accordingly, purified LT-α1/B2 complexes bind with high affinity to LT-B-R between adjacent LT-B subunits, but lack α/α clefts and thus do not cross-activate signaling through the TNF receptors. Thus the LT- α/β heteromeric 20 complexes of this invention will not stimulate TNFassociated inflammatory responses.

LT- α 1/ β 2 administration does not activate endothelial cell changes associated with inflammatory response even at relatively high levels. For this reason, the side effects due to activation of the inflammatory cascades observed with TNF should not be a problem using the pharmaceutical compositions and treatment methods to activate the LT- β -R.

Human LT- α 1/ β 2 binds to mouse LT- β -R essentially as well as to human LT- β -R. Injection of 100 µg human LT- α 1/ β 2 per mouse is not lethal (Example 11), suggesting that stimulation of LT- β -R in the whole animal

does not hav the overt toxicity seen when similar experiments were tried with FasR or p60 TNF-R activation.

The use of specific anti-LT-ß-R monoclonal antibodies or antibody combinations to trigger this pathway in humans may have several advantages over treatment with LT- α/β heteromeric complexes. An anti-receptor antibody therapy will be more selective than treating with ligand. Moreover, recombinant forms of anti-LT-ß-R mAbs would be easier to engineer and produce in large scale than the soluble LT- α/β heteromeric complexes.

5

10

15

20

25

30

It is envisioned that the mAbs or LT- α/β heteromeric complexes would be administered to tumorbearing people in conjunction with a conventional antitumor therapy (i.e. radiation and chemotherapy). A combined treatment of LT- β -R activation with conventional chemotherapies may provide an extra factor of tumor killing activity that would be more likely to clear a patient of tumorigenic cells than when conventional antitumor therapy is used alone.

It is further possible that this approach may have relatively few side effects and therefore could be given in a semi-prophylactic sense in cases of carcinomas that may not have metastasized, or in patients from families who show a genetic pre-disposition for a certain type of cancer.

The following are examples which illustrate the LT- α/β heteromeric complexes and the anti-LT- β -R mAbs of this invention and the methods used to characterize them. These examples should not be construed as limiting: the examples are included for purposes of illustration and the present invention is limited only by the claims.

- 39 -

EXAMPLE 1

Generation f Baculovirus-infected Insect Cell Supernatants Containing $LT-\alpha/\beta$ Forms

Recombinant baculovirus encoding either full length LT- α or a secreted form of LT- β were made as 5 described (Crowe et al., Science, 264, pp. 707-710 (1994)). High five insect cells (Invitrogen, San Diego, CA.) were inoculated at a density of 2x10⁵ cells/ml into 7.2 liters of SF 900-II (Gibco) media without serum. culture reached 1.8x106 cells/ml 48 hours later and was 10 infected with 150 ml (3x108 PFU/ml) of LT-B and 300 ml of $LT-\alpha$ baculovirus stocks. Two days later, the culture was harvested and the cell debris was removed by centrifugation. After addition of EDTA and PMSF (1 mM EDTA and 150 µM PMSF final concentration), the clarified 15 supernatant was concentrated 10-fold by ultrafiltration using a S1YM10 (Amicon) spiral cartridge. The concentrate was divided into six 120 ml portions and aliquots were stored at -70 C prior to purification.

20

EXAMPLE 2

Preparation of Soluble LT-8 Receptors as Immunoglobulin Fc Chimera

The extracellular domain of LT-B-R up to the transmembrane region was amplified by PCR from a cDNA clone using primers that incorporated NotI and SalI 25 restriction enzyme sites on the 5' and 3' ends, respectively (Browning et al., J. Immunol., 154, pp. 33-46 (1995)). The amplified product was cut with NotI and SalI, purified and ligated into NotI-linearized vector pMDR901 along with a SalI-NotI fragment encoding the Fc 30 region of human IgG1. The resultant vector contained the dihydrofolate reductase gene and the LT-B-R-Fc chimera driven by separate promoters. The vector was electroporated into CHO dhfr cells and methotrexateresistant clones were isolated as per standard 35

- 40 -

procedures. The LT-B-R-Fc is secreted into the medium and an ELISA assay was used to select for cell lines producing the highest level of the chimeric protein. A high-producing cell line was grown to large numbers and conditioned medium collected. The pure protein was isolated by Protein A sepharose fast flow affinity chromatography.

5

10

15

20

25

EXAMPLE 3

Affinity Purification of LT-α1/82 Using TNF-R and LT-8-R

To prepare resins for the receptor affinity purification of LT forms, purified preparations of LT-B-R-Fc (as described in Example 2 herein) and TNF-R p60-Fc (Crowe et al., <u>Science</u>, 264, pp. 707-10 (1994)) were immobilized on CNBr-sepharose (Pharmacia) at 5 mg/ml resin essentially following the manufacturer's specifications. The resins were put through one elution cycle prior to use. A portion (120 ml) of the S1Y10 concentrate was passed over two sequential p60 TNF-R-Fc columns, which bind LT- α and LT- α 2/B1. The flow through, which contained LT-α1/β2 and LT-β, was passed over a LT-B-R-Fc column. The column was washed with 5 volumes each of PBS, PBS with 0.5 M NaCl and PBS, and then the LT- α and LT- α 2/ β 1 complexes were eluted with 25 mM sodium phosphate, 100 mM NaCl, pH 3.5. Elution fractions were immediately neutralized with 1/20 volume of 0.5 M sodium

phosphate, pH 8.6 and stored on ice. Fractions containing protein were identified by absorbance at 280 nm, peak fractions were pooled and the elution pools from the columns were analyzed by SDS-PAGE stained with coomassie brilliant blue. Elution as described above yielded greater than 95% pure LT-α1/β2.

5

10

15

20

25

30

EXAMPLE 4

Characterization of the Purified LT- α 1/82 Ligands

The fractions of Example 3 were sized by gel exclusion chromatography to assess whether trimers were formed and if aggregates were present. A TSK G3000 sw x2column was used at a flow rate of 0.5 ml/min to size separate a BioRad gel filtration protein standard, and the three different LT trimers, LT- α 3, LT- α 2/ β 1 and LT α 1/ β 2. Figure 1A shows that very little, if any of the LT- α 1/ β 2 trimers show up as high molecular weight aggregate. Comparison to size standards shows that the three forms are all trimeric, i.e. about 50-60 kDa. Assuming the trimer, the stoichiometry of LT- α to LT- β contained in the purified LT- α 1/ β 2 and LT- α 2/ β 1 fractions was evaluated by either densitometry of the coomassie stained gels or by peak height analysis of the two peaks following resolution on C4 reverse phase HPLC. Both measurements confirmed the identity of the fractions eluted off the affinity columns as described above.

The purity of the preparations was further assessed by ion exchange chromatography using BioCAD instrumentation to run pH maps of LT- α 1/ β 2 and LT- α 2/ β 1 on the weak cation exchanging resin under several different buffer systems. The method that exhibited the greatest ability to cleanly retain and separate the three trimers incorporated a POROS CM (carboxymethyl) column run at 5 ml/min with a 16.66 mM MES, 16.66 mM HEPES, 16.66 mM Na Acetate pH 6.5 buffer and eluting with a 1 M NaCl gradient over 20 column volumes. The BioCAD chromatograms of LT- α 1/ β 2 and LT- α 2/ β 1 complexes are shown in Figure 1B. Each trimer, LT- α 3, LT- α 2/ β 1 and LT- α 1/ β 2 eluted at a different salt concentration and there was no evidence for cross contamination of more than 1-2% in the various preparations.

25

30

- 42 -

EXAMPLE 5

Killing of HT29 Human Adenocarcinoma Cells by Soluble LT-α1/B2 Complexes

The HT29 cytolytic assay has been previously described (Browning and Ribolini, J. Immunol., 143, pp. 5 1859-67 (1989)). In a typical assay, serial dilutions of LT- α 1/ β 2 (and other cytokines where applicable) were prepared in 0.05 ml in 96 well plates and 5000 trypsinized HT29-14 cells added in 0.05 ml of media containing 0 or 80 U/ml (anti-viral units) of human 10 IFN-y. HT29-14 cells are from a subclone of the original ATCC-derived HT29 line that is more homogeneous. HT29-14 cells were used in the assays; all of these results can also be observed using the original ATCC-derived HT29 line. After 3-4 days, mitochondrial reduction of the dye 15 MTT was measured as follows: 10 l of MTT was added and after 3 hours, the reduced dye dissolved with 0.09 ml of isopropanol with 10mM HCl, and the O.D. measured at 550 nm. Soluble receptor forms prepared as described herein, or pure human IgG were added in 10 µl prior to 20 the cells to give a final concentration of 5 µg/ml.

Figure 2A shows the killing of HT29 cells by treating with anti-Fas receptor mAb CH-11 (which stimulates FasR signaling); TNF, LT- α 3, LT- α 1/ β 2 and LT- α 2/ β 1 ligands in conjunction with IFN- γ . Visual inspection of the cells treated with LT- α 1/ β 2 reveals that this agent kills cells rather than just blocking cell proliferation. In the absence of IFN- γ , no effects are observed, reflecting the unusual ability of IFN- γ to influence how cells interpret signaling from the TNF family of receptors. Interferons α and β were 100-fold less effective than IFN- γ as quantitated based on antiviral activity units.

Figur 2B shows the inhibition of LT- α 1/ β 2
35 killing by soluble LT- β -R-Fc but not p60-TNF-R-Fc,

- 43 -

demonstrating that cytotoxicity is specific to LT- α 1/ β 2. The lack of inhibition by p60-TNF-R-Fc indicates that contaminating LT- α (known to be less than 1%) cannot account for the cytotoxic activity of LT- α 1/ β 2.

5

EXAMPLE 6

Anti-LT-8-R mAbs Potentiate the Killing of HT29 Cells by LT- α 1/82 Complexes

Cytolytic assays were performed as described in Example 5, except that IFN-y and anti-LT-B-R mAbs (0.01 -1000 ng/ml series) were added to the cells at 2x final 10 concentration and then 50 µl of the cell solution were added to the wells containing diluted LT- α 1/ β 2. Growth was assessed as described in Example 5. Figure 3 shows the differential effects of two different anti-LT-B-R mAbs in their ability to potentiate LT- α 1/ β 2 cytotoxic 15 activity. Figure 3A shows that the anti-LT-B-R mAb CDH10 potentiates LT- α 1/ β 2 cytotoxic activity in a dosage-dependent manner. Figure 3B shows the effects of another anti-LT-B-R mAb, BDA8, in the same assay. The BDA8 mAb inhibits the cytotoxic activity of LT- α 1/ β 2 20 rather than potentiating tumor cell death.

EXAMPLE 7

Immobilized Anti-LT-8-R mAbs Can Kill HT29 Tumor Cells

To immobilize anti-LT-ß-R mAbs onto a plastic surface, 96-well tissue culture plates were coated with 50 µl of 10 µg/ml goat anti-mouse Fc polyclonal antibody (Jackson ImmunoResearch), washed and blocked with 5% FCS in PBS, followed by capture of the indicated anti
LT-ß-R mAb and another wash. HT29 cells were plated into the mAb-coated wells, and cytolytic assays were conducted

- 44 -

as in Example 5. Figure 4A illustrates the cytotoxic effects of immobilized BDA8 and CDH10 anti-LT-ß-R mAbs on HT29 cells. Each mAb individually elicits cytotoxicity on tumor cells when it is immobilized onto a surface.

Figure 4B shows that the same BDA8 and CDH10 anti-LT-ß-R mAbs tested individually in solution are not cytotoxic and thus the cytolytic activity of a single anti-LT-ß-R mAb in vitro appears to be a function of its immobilization.

10 EXAMPLE 8

5

15

20

A combination of anti-LT-8-R mAbs in solution directed against distinct epitopes kill HT29 cells

Growth of HT29 cells was assessed as described in Example 5 except that either one or two anti-LT-\(\beta\)-R mAbs were included in the growth medium. Table 1 shows the effects on HT29 cells observed when various anti-LT-\(\beta\)-R mAbs were included in solution (i.e., not immobilized on plastic). The anti-LT-\(\beta\)-R mAbs can be arranged into groups I-IV based on their relative abilities to work in combination with each other in a HT29 cytolytic assay. The anti-LT-\(\beta\)-R mAbs results generated in cytolytic assays parallel receptor binding data which suggest that the mAbs in each different group recognize different epitopes of the LT-\(\beta\)-R.

Table 1. Combinations of soluble anti-LT-β-R mAbs are cytotoxic to human adenocarcinoma HT29 cells. Anti-LT-β-R mAbs are grouped into Groups I, II, III and IV based on their effects in combination with each other in HT29 cell cytolytic assays. Pluses refer to the relative level of cytolytic effects of the mAb combination on HT29 cells in the presence of 80U/ml IFN-γ. nr = not relevant; nd = not determined.

						Second	mAb		
10	Group	First _mAb	Group BDA8	I AGH1		II BHA10			Group IV CBE11
	I	BDA8 AGH1	nr -	- nr	+ ++	++ +++	+ ++	nd nd	nd nd
15	II	BCG6 BHA10	++ ++	++	nr + -	- nr	+++	+ nd +++	nd ++++
	III	BKA11 CDH10	+ ++	++ ++	+++	nd +++	nr -	- nr	nd +++
	IV	CBE11	nd	+	+	+++-	+ nd	+++	+ nr

Figure 5A-D quantitates the effects of including in the

HT29 cytolytic assay representative pairwise combinations of cooperating anti-LT-B-R mAbs directed against different epitopes of LT-B-R. Figure 5A shows the cytotoxic effects of BHA10 and CBE11, Figure 5B of CDH10 and CBE11, and Figure 5C of CDH10 and AGH1, alone and in combination. Figure 5D shows the cytotoxic effects of the CDH10 and AGH1 mAb combination in a different tumor cell line called WiDr.

Table 2 summarizes the characteristics of the representative anti-LT-B-R mAbs of this invention.

Table 2

Summary of Mouse Anti-human LT-6-R mAbs

- 46 -

P
-
ū
X.
즼
9
3
٦
1
Ŋ
-

ë Vith	100 EC		iates	ğ	777
Soluble mAb with Ltalll	inhibits inhibits	mixed	potentiates potentiates	no effect	no effect no effect no effect
Soluble mAb alone	P-/+	+ +	· +	-/+	1 1 1
mAb Immobilized on Plastic	+	+ +	+ +	+	1 1 1
Blocking Receptor Binding ^b	‡‡	‡ ‡	<u>+</u> +	‡	- pu
Cell Staining	‡‡	‡‡	‡‡	‡	ı ı p ı
mAb Name	BDA8 AGHI	BCG6 BHA10	BKA11 CDH10	CBEII	MOPC21 HT29/26 TS 2/9°
mAb Group		=	HH	N	Controls

FACS staining of CHO cells transfected with the LT-β-R.

bAssay assessed whether antibody blocks binding of soluble receptor to the activated T-cell hybridoma II-23. nd = not done. CHT-29 cells were grown with IFN-γ on anti-LT-β-R coated plates as defined in the methods.

Variable, partial inhibition in some assays, no effects in others.

Anti-human LFA-3, a mouse IgG1.

25

30

- 47 -

EXAMPLE 9

Reliance on Endogenous IFN- γ For the Treatment of Tumor Cells

IFN-y, a preferred LT-B-R activating agent of 5 the present invention, is a cytokine which exhibits antitumor activity and which is tolerated in humans. Endogenous IFN-y present in the environment surrounding a tumor may be at sufficiently high concentrations to function as a LT-B-R activating agent of this invention without adding exogenous IFN-y. The concentration of 10 IFN-y in the vicinity of a tumor may be ascertained using standard immunochemical techniques with tissue samples from the region of the tumor. If the endogenous concentration of IFN-y is high enough to elicit anti-15 tumor activity in combination with the LT-α/β heteromeric complexes or anti-LT-B-R mAbs of this invention (as determined by the cytolytic assays described herein), then IFN-y need not be administered as a second LT-B-R activating agent in the compositions or methods of this 20 invention.

EXAMPLE 10

Induction of Endogenous IFN-y As A LT-8-R Activating Agent For the Treatment of Tumor Cells

Compounds which can induce the endogenous production of interferons such as IFN-y fall within the group of LT-ß-R activating agents of this invention. For example, interferons can be induced by treating with double-stranded RNA molecules such as polyriboguanylic/polyribocytidylic acid (poly-G/C).

Female C57/b16 (6-8 weeks old) can be injected with 18 mg (600 mg/kg) of D-galactosamine which sensitizes mice to the effects of TNF and other antitumor agents. A series of concentrations of poly-G/C (Juraskova et al., Eur. J. Pharmacol., 221, pp. 107-11

- 48 -

(1992)) in a neutral saline solution is added to purified LT- α 1/ β 2 (10-100 μ g) and the solution administered to mice as an intraperitoneal (i.p.) injection. The anti-tumor activity of LT- α 1/ β 2 will be enhanced by the presence of poly-rG/rC double stranded RNA.

Similarly, the interferon stimulator from the plant Glycyrrhiza glabra (Acharya et al., Indian J. Med. Res., 98, pp. 69-74 (1993)) may be administered to humans intravenously at doses ranging from 40-100 ml/day. The optimal dose for LT- β -R activation in the presence of either LT- α/β heteromeric complexes or anti-LT- β -R Abs may be determined empirically and will depend on factors such as the tumor type, mode of delivery and delivery schedule.

10

Imiquimod R-837 (Bernstein et al., Antiviral Res., 20, pp. 45-55 (1994)); Saparal (Paramonova et al., Yopr. Virusol., 39, pp. 131-34 (1994)); Bropirimine (Onishi and Machida, Hinyokika Kiyo, 40, pp. 195-200 (1994)); or Ridostin (Cheknev et al., Yopr. Virusol., 39, pp. 125-28 (1994)), may also be administered as LT-β-R activating agents in conjunction with LT-α/β heteromeric complexes, anti-LT-β-R Abs, or a combination thereof. In each case, the preferred modes of delivery and optimal doses can be determined empirically using the published reports as starting points for optimization by routine clinical procedures.

EXAMPLE 11

Mice Tolerate Injections of Human LT- α 1/ β 2

Female C57/b16 (6-8 weeks old) acclimated to

the facility for several days were injected i.p. with 18

mg (600 mg/kg) of D-galactosamine, which sensitizes mice

to the effects of TNF and other anti-tumor agents.

Either human TNF, LT-α or LT-α1/β2 was then given i.p.

- 49 -

Table 3 documents the survival of treated mice 24 hours after injection.

Table 3

5	<u>Agent</u>	Dose (ug/animal)	Survival
	saline	-	4/4
	hu-TNF	0.2	0/6
	hu-TNF	1.0	0/2
	hu-TNF	10	0/4
10	hu-LT-α	0.2	2/2
	hu-LT-α	1.0	2/2
	hu-LT-α1/B2	10	2/2
	$hu-LT-\alpha1/\beta2$	100	2/2

15

20

25

EXAMPLE 12

Construction of a recombinant anti-LT-8-R IgM monoclonal antibody

Using the anti-tumor cytotoxicity assays described above coupled with standard tumor growth models in immunodeficient mice, an anti-LT-B-R IgG with suitable properties can be selected. Universal primers which hybridize to each of the variable domains of the IgG heavy and light chains of the selected anti-LT-B-R IgG can be used to prepare variable domain DNA from RNA isolated from the secreting hybridoma cell line using standard reverse transcriptase/PCR methodologies. These protocols have been described (Arulanandam et al., J. Exp. Med., 177, pp. 1439-50 (1993); Lane et al., Eur. J. Immunol., 22, pp. 2573-78 (1993); Traunecker et al., Nature, 339, pp. 68-70 (1989)).

30 The amplified products are then assembled into vectors containing the human CH1, CH2 and CH3 μ chain domains. Co-expression of the two chains in a single host will allow assembly of the heavy and light chains into a pentameric IgM molecule. This molecule is a

chimera composed of mouse variable regions coupled to human constant regions.

5

10

25

30

Alternatively, a process using PCR to amplify DNA encoding only the actual binding regions of the variable regions can be used. Amplified DNA is then inserted into vectors containing all of the human IgG sequences except for the actual amino acids involved in binding the antigen. Such constructs are called "humanized" antibodies and the detailed methods for their production are well-known (e.g. WO 94/04679).

EXAMPLE 13

Anti-LT-8-R IgM Monoclonal Antibodies Function as LT-8-R Activating Agents

Anti-LT-B-R IgM antibodies can be prepared in a recombinant form as described in Example 12.

Alternatively, complete mouse IgMs isolated by hybridoma fusion techniques using primary immunization of normal mice or extensive immunization of CD40 signaling-deficient mice (Kawabe et al., Immunity, 1, pp. 167-78

(1994); Xu et al., Immunity, 1, pp. 423-31 (1994)) can be used as a source of anti-LT-B-R IgM mAbs.

Anti-LT- β -R IgM mAbs will be significantly more potent as LT- β -R activating agents than their normal bivalent IgG counterparts as measured by dose-response comparisons in the HT29 cytolytic assay in the presence of IFN- γ . The anti-LT- β -R IgM mAbs function as LT- β -R activating agents both when they are immobilized and when they are administered in solution. In addition, we expect that they will augment the anti-tumor activity of LT- α/β heteromeric complexes.

25

30

35

EXAMPLE 14

Anti-LT-8-R Monoclonal Antibodies Inhibit the Growth of Human Tumor Cells in SCID Mice

Balb/c SCID mice (Jackson Labs, Bar Harbor, ME) 5 were injected with 1 x 106 trypsinized and washed human adenocarcinoma WiDr cells in a volume of 0.2 ml of PBS subcutaneously (s.c.) onto the back of the animal. Injected WiDr cells form tumors in the mice, and the 10 ability of an anti-LT-B-R mAb to inhibit tumor growth was monitored. In one set of experiments, mice were treated with or without the CBE11 anti-LT-B-R mAb -- either with or without human IFN-y (106 antiviral units/mouse) -- at the same time as the WiDr cells were inoculated s.c. 15 (Figure 6A). Antibodies and IFN-y were administered alone or together by i.p. injection in 0.2 ml. Control mice were injected with saline alone, IFN-y alone, or a control anti-human LFA-3 mAb (1E6) with IFN-y. The size of each resulting tumor was scored 30 days after 20 inoculation. Tumor volume (in cc) was calculated from the radius as determined by caliper measurements in two dimensions. Animals treated with CBE11 or 1E6 mAbs received 10 µg/mouse or 50 µg/mouse of antibody (Figure 6A; circles with dots and open circles, respectively).

In another set of experiments, mice were inoculated s.c. with the WiDr cells and tumors were allowed to grow for 15 days before the mice were treated with the CBE11 anti-LT-B-R mAb (Figure 6B). At day 15 (before antibody treatment), the average tumor volume was 0.076 cc with an average diameter of 0.53 cm. The CBE11 anti-LT-B-R mAb (50 µg) was then administered — either with or without human IFN-y (106 antiviral units/mouse) — by i.p. injection in 0.2 ml to a group of 12 animals. Injections were repeated three more times over a three week period. Control groups (12 mice/group) were injected with IFN-y alone (106 antiviral units/mouse) or

with 50 µg of a control anti-human LFA-3 mAb (1E6) + IFN-Y (106 antiviral units/mouse). The growth of the tumors present at day 15 was scored over time, from 15 to 49 days post-tumor cell inoculation. The results shown in Figure 6B were determined in a blinded format. Tumors treated with CBE11 mAb either with or without IFN-Y stopped growing. Following three injections of CBE11 mAb (+/- IFN-Y) over three weeks, tumor growth was arrested for at least 7 weeks post-inoculum, at which time the experiment was terminated.

10

Applicants or agents file			International application No.
	B185 CI	ው ውሮጥ	
reference number	DIOJ CI	FFCI	

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made below relate to the microorganism re	elemed to in the description
on page 28, line S _18	
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet X
Name of depositary institution	
American Type Culture Coll	ection
Address of depositary institution (including posici code and count	(4)
12301 Parklawn Drive Rockville, Maryland 20852 United States of America	
Identification Reference by De	epositor: Hybridoma, CB.Ell.l
Date of deposit	Accession Number
12 January 1995 (12.01.95	НВ 11793
C. ADDITIONAL INDICATIONS (Icome plant of non applica	This information is continued on an additional about
posited microorganisms will be lication of the mention of the until the date on which the apportise deemed to be withdrawn, a Implementing Regulations under sample to an expert nominated by	grant of the European patent or plication is refused or withdrawn as provided in Rule 28(3) of the the EPC only by the issue of a
E. SEPARATE FURNISHING OF INDICATIONS (Id	rve blank if not applicable)
The indications listed below will be submitted to the Internation Number of Deposit?)	
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	This sheet was received by the international Bureau on:
Authorized officer	Authorized officer
,	

				International application No.
Applicants or agents tile reterence number	B185	CIP	PCT	International

in the description
A. The indications made below relate to the microorganism referred to in the description on page 28 line S 18-26
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet X
Name of depositary institution
American Type Culture Collection
Address of depositary institution including postal code and country)
12301 Parklawn Drive Rockville, Maryland 20852 United States of America
Identification Reference by Depositor: Hybridoma, BC.G6.AF5
Date of deposit Accession Number
12 January 1995 (12.01.95) HB 11794
C. ADDITIONAL INDICATIONS (Icave plank if not applicable) This information is continued on an additional about
In respect of the designation of the EPO, samples of the deposited microorganisms will be made available until the publication of the mention of the grant of the European patent or until the date on which the application is refused or withdrawn or is deemed to be withdrawn, as provided in Rule 28(3) of the Implementing Regulations under the EPC only by the issue of a sample to an expert nominated by requester (Rule 20(4) EPC). D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (i) the indicators are not for all designed States) EPO
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)
The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications e.g., "Accessed Number of Deposit")
For receiving Office use only
This sheet was received with the international application This sheet was received by the International Bureau on:
Authorized officer Authorized officer

Applicants or agents file B185 CIP PCT	 			
reference number B103 CIF FCI	B185	CIP	PCT	International application No.

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made below relate to the microorganism retern on page 28 time S 18-	erred to in the description 26
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution	
American Type Culture Colle	ction
Address ut depositary institution tractading poster code and country)	
12301 Parklawn Drive Rockville, Maryland 20852 United States of America	
Identification Reference by Dep	positor: Hybridoma, BH.Al0
Date of deposit 12 January 1995 (12.01.95)	Accession Number HB 11795
C. ADDITIONAL INDICATIONS (Icave blank if not applicable	this information is continued on an additional about
In respect of the designation posited microorganisms will be a lication of the mention of the cuntil the date on which the apport is deemed to be withdrawn, as Implementing Regulations under sample to an expert nominated by D. DESIGNATED STATES FOR WHICH INDICATION	grant of the European patent or lication is refused or withdrawn s provided in Rule 28(3) of the the EPC only by the issue of a requester (Rule 28(4) EPC)
E. SEPARATE FURNISHING OF INDICATIONS (leave	blank if not applicable)
The indications listed below will be submitted to the International Number of Deposit 7	Buresu later (specify the general nature of the indications a.g., "Accumen
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	This sheet was received by the international Bureau on:
Authorized officer	Authorized officer

				International application No.
Applicants or agents file reference number	B185	CIP	PCT	

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made below relate to the microorganism referred to in the description on page 28 , line S 18-26
was desirable and distinguished V
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional speece [A.]
Name of depositary institution
American Type Culture Collection
Address of depositary institution tinetimine postal code and country)
12301 Parklawn Drive
Rockville, Maryland 20852 United States of America
Identification Reference by Depositor: Hybridoma, AG.Hl.5.1
Date of deposit Accession Number
12 January 1995 (12.01.95) HB 11796
C. ADDITIONAL INDICATIONS (leave nlank if not applicable) This information is continued on an additional about
In respect of the designation of the EPO, samples of the deposited microorganisms will be made available until the publication of the mention of the grant of the European patent or until the date on which the application is refused or withdrawn or is deemed to be withdrawn, as provided in Rule 28(3) of the Implementing Regulations under the EPC only by the issue of a sample to an expert nominated by requester (Rule 26(4) EPC). D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (6) the indicators are not for all designed States) EPO
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)
The indications listed below will be submitted to the international Bureau later (specify the general nature of the indications a.g., "Accommon Number of Depart")
For receiving Office use only
This about was received with the international application This about was received by the International Bureau on:
Authorized officer Authorized officer

Applicants or agents file				International application No.
Applicants or agents file reference number	B185	CIP	PCT	(,
TOTOTON NOTICE				

A. The indications made below relate to the microorganism referred to in the description on page 28 line S 18-26					
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional abeet					
Name of depositary institution					
American Type Culture Collection					
Address of depositary institution (including poste) code and country)					
12301 Parklawn Drive Rockville, Maryland 20852 United States of America					
Identification Reference by Depositor: Hybridoma, CD.H10.1					
Date of deposit 12 January 1995 (12.01.95) Accession Number HB 11797					
C. ADDITIONAL INDICATIONS ((core blank if not applicable) This information is continued on an additional about					
In respect of the designation of the EPO, samples of the deposited microorganisms will be made available until the publication of the mention of the grant of the European patent or until the date on which the application is refused or withdrawn or is deemed to be withdrawn, as provided in Rule 28(3) of the Implementing Regulations under the EPC only by the issue of a sample to an expert nominated by requester (Rule 28(4) EPC). D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (1) the indications are as for all designed State) EPO					
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)					
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications a.g., "Accustmen Number of Depart")					
For receiving Office use only For laternational Bureau use only					
This sheet was received with the international application This sheet was received by the International Bureau on:					
Authorized officer Authorized officer					

			International application No.
Applicants or agent's tile	B185 CIP	PCT	interimental and a second
reference number	2200 0		

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made believe relate to the microorganism refer	rred to in the description 26						
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet						
Name of depositary institution							
American Type Culture Colle	ction						
Address of depositary institution including poster code and country)							
12301 Parklawn Drive Rockville, Maryland 20852 United States of America	Rockville, Maryland 20852						
Identification Reference by Der							
Date of deposit 12 January 1995 (12.01.95)	Accession Number HB 11798						
C. ADDITIONAL INDICATIONS (leave plank if not applicable	This information is continued on an additional about						
In respect of the designation of the EPO, samples of the deposited microorganisms will be made available until the publication of the mention of the grant of the European patent or until the date on which the application is refused or withdrawn or is deemed to be withdrawn, as provided in Rule 28(3) of the Implementing Regulations under the EPC only by the issue of a sample to an expert nominated by requester (Rule 2%(4) EPC). D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designed States) EPO E. SEPARATE FURNISHING OF INDICATIONS (have blank if not applicable) The indications insted below will be submitted to the international Bureau later (specify the general nature of the indications as a. "Accounter.")							
Number of Depart*)							
For receiving Office use only	For International Bureau use only						
This sheet was received with the international application	This sheet was received by the interastional Bureau on:						
Asthonized officer	Authorized officer						

				No.
Applicants or agents file				International application No.
	B185	CTP	PCT	
reference number	2103			

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made below relate to the microorganism referred to in the description on page 28 line S 18-26						
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet						
Name of depositary institution						
American Type Culture Collection						
Address at depositary institution tinetuding postel code and country)						
12301 Parklawn Drive Rockville, Maryland 20852 United States of America						
Identification Reference by Depositor: Hybridoma, BK.All.ACLC						
12 January 1995 (12.01.95) Accession Number HB 11799						
C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional about						
In respect of the designation of the EPO, samples of the deposited microorganisms will be made available until the publication of the mention of the grant of the European patent or until the date on which the application is refused or withdrawn or is deemed to be withdrawn, as provided in Rule 28(3) of the Implementing Regulations under the EPC only by the issue of a sample to an expert nominated by requester (Rule 20(4) EPC). D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designed State) EPO						
E. SEPARATE FURNISHING OF INDICATIONS (lueve blank if not applicable)						
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications a.g., 'Accommon Number of Deposit')						
For receiving Office use only For International Buresu use only						
This sheet was received with the international application This sheet was received by the International Bureau on:						
Authorized officer Authorized officer						

A				International application No.
Applicant's or agent's tile	B185	CID	DCT	1
reference number	DIOD	CIP	FCI	}

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made below relate to the microorganism reterred to in the description on page 28 line S 18-26					
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet X					
Name of depositary institution American Type Culture Collection					
Address in depositary institution (including postal code and country)					
12301 Parklawn Drive Rockville, Maryland 20852 United States of America					
Identification Reference by Depositor: Hybridoma, CB.Ell.1					
Date of deposit 12 January 1995 (12.01.95) Accession Number HB 11793					
C. ADDITIONAL INDICATIONS (Icave plank if not applicable) This information is continued on an additional about					
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.					
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated Status)					
Finland					
E. SEPARATE FURNISHING OF INDICATIONS (Icave blank if not applicable)					
The indications listed below will be submitted to the International Bureau later (spacify the general nature of the indications e.g., "Accommon Number of Deposit")					
For international Bureau use only					
This sheet was received with the international application This sheet was received by the International Bureau on:					
Authorized officer Authorized officer					

Applicant's or agent's file				International application No.	
reference number	B185	CIP	PCT		

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made nellow relate to the microorganism on page 28 , line S 1						
100 100 100 100 100 100 100 100 100 100						
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional abeet					
Name of depositary institution						
American Type Culture Co	llection					
Address of depositary institution fincing postal code and coun	LLÝI					
12301 Parklawn Drive						
Rockville, Maryland 20852 United States of America	2					
	Depositor: Hybridoma, BC.G6.AF5					
Date of deposit	Accession Number					
12 January 1995 (12.01.95)) HB 11794					
C. ADDITIONAL INDICATIONS (Icave plant if not applied	cable) This information is continued on an additional sheet					
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.						
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)						
Finland						
E. SEPARATE FURNISHING OF INDICATIONS (A						
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications a.g., "Accusion Number of Depast")						
For receiving Office use only	For International Bureau use only					
This sheet was received with the international application	11 <u> </u>					
Authorized officer	Authorized officer					

						-
ı	Application and account of the				International application No.	
	Applicants or agents tile				1 '''	
i	reference number	B185	CIP	PCT_{-}		

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made negow relate to the microorganism rete on page 28 , line S 18-	-26						
on page, line							
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet X						
Name of depositary institution							
American Type Culture Collection							
Address ist depositary institution tineimine postal code and country)							
12301 Parklawn Drive							
Rockville, Maryland 20852							
United States of America							
Identification Reference by Dep							
Date of deposit	Accession Number HB 11795						
12 January 1995 (12.01.95)	нв 11795						
C. ADDITIONAL INDICATIONS (leave alank if not applicable	This information is continued on an additional sheet						
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.							
D. DESIGNATED STATES FOR WHICH INDICATION	NS ARE MADE (if the indications are not for all designated States)						
Finland							
E. SEPARATE FURNISHING OF INDICATIONS (Icon	e blank if not applicable)						
The indications listed below will be submitted to the International							
Number of Deposit")							
For receiving Office use only	For International Bureau use only						
This abeat was received with the international application	This abcet was received by the International Bureau on:						
Authorized officer	Authorized officer						
remained nitre							

5	2	1	1 1	
	_ ,	٠.		

Applicant's or agent's file				International application No.
reference number	B185	CIP P	CT	

A. The indications made nellow relate to the microorganism reterred to in the description	
on page28line s 18-26	
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional abeet	X
Name of depositary institution	_
American Type Culture Collection	
Address of depositary institution finetiming poster code and country)	
12301 Parklawn Drive	
Rockville, Maryland 20852	
United States of America	
Identification Reference by Depositor: Hybridoma, AG.Hl.5.	1
Date of deposit 12 January 1995 (12.01.95) Accession Number HB 11796	
C. ADDITIONAL INDICATIONS (Icave plant if not applicable) This information is continued on an additional about	X
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.	
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designment States)	(23)
Finland	
E. SEPARATE FURNISHING OF INDICATIONS (Icere blank if not applicable)	
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Account Number of Deposit")	101
For receiving Office use only For International Bureau use only	
This sheet was received with the international application This sheet was received by the international Bureau of	30:
Authorized officer Authorized officer	ᅱ
	-

			International application No.	
Applicants or agent's tile reference number	B185 CIP	PCT		

eterreit to in the description
. The indications made decide relate to the microorganism referred to in the description
3. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet X
Name of depositary institution
American Type Culture Collection
Address of depositary institution fineitiding postal code and country)
12301 Parklawn Drive Rockville, Maryland 20852 United States of America
Identification Reference by Depositor: Hybridoma, CD.H10.1
12 January 1995 (12.01.95) HB 11797
C. ADDITIONAL INDICATIONS (leave alank if not applicable) This information is continued on an additional about
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (i) the indications are not for all designment Status)
Finland
E. SEPARATE FURNISHING OF INDICATIONS (lusve blank if not applicable)
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., 'Accession Number of Deposit')
O company was called
For receiving Office use only This sheet was received with the international application This sheet was received by the International Bureau on
Authorized officer Authorized officer

Amelianasa an anasasa at la				International application No.
Applicant's or agent's tile	D10E	OTD	nom.	
reference number	B185	CIP	PCT	· ·
. otal olive il minde.				

A. The indications made nellow relate to the microorganism reterred to in the description on page 28 line S_18-26
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional about X
Name of depositary institution American Type Culture Collection
Address of depositary institution fineithning poster code and country)
12301 Parklawn Drive Rockville, Maryland 20852 United States of America Identification Reference by Depositor: Hybridoma, BD.A8.AB9
Date of deposit 12 January 1995 (12.01.95) Accession Number HB 11798
C. ADDITIONAL INDICATIONS (Icave plant if not applicable) This information is continued on an additional about
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)
Finland
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications a.g., "Accesses Number of Depart")
For receiving Office use only
This sheet was received with the international application
Authorized officer Authorized officer

	52/14	
Applicant's or agent's file reference number	B185 CIP PCT	International application No.

A. The indications made nellow relate to the microorganism referred to in the description on page 28 , line S 18-26
on page 28 line S 18-26
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional abest
Name of depositary institution
American Type Culture Collection
Address of depositary institution tractituing postal code and country)
12301 Parklawn Drive Rockville, Maryland 20852 United States of America
Identification Reference by Depositor: Hybridoma. BK.All.AC
Date of deposit Accession Number
12 January 1995 (12.01.95) HB 11799
C. ADDITIONAL INDICATIONS (Iceve plent if not applicable) This information is continued on an additional sheet
In respect of the designation of Finland, until the application has been laid open to public inspection by the Finnish Patent Office, or has been finally decided upon by the Finnish Patent Office without having been laid open to public inspection, samples of the deposited microorganisms will be made available only to an expert in the art.
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated State
Finland
E. SEPARATE FURNISHING OF INDICATIONS (losve blank if not applicable)
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Access Number of Deposit")
For receiving Office use only
This sheet was received with the international application This sheet was received by the International Bureau
Authorized officer Authorized officer

_	2	•	1	_
כ	Z	/	1	2

	32/	15	
Applicants or agent's tile reference number	B185 CIP	PCT	International application No.

A. The indications made nellow relate to the microorganism referred to in the description on page
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional sheet X
Name of depositary institution American Type Culture Collection
Address of depositary institution inclining posial code and country
12301 Parklawn Drive Rockville, Maryland 20852 United States of America
Identification Reference by Depositor: Hybridoma, CB.Ell.l
12 January 1995 (12.01.95) Accession Number HB 11793
C. ADDITIONAL INDICATIONS (leave nlank if not applicable) This information is continued on an additional about
Applicant(s) hereby give notice of my/our intention that samples of the above-identified culture shall be available only to experts in accordance with paragraph 3 of the Fourth Schedule to the Patents Rules 1995.
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designment Scane)
Singapore
E. SEPARATE FURNISHING OF INDICATIONS (Icone blank if not applicable)
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications a.g., 'Accessed Number of Deposes')
For International Bureau use only
This sheet was received with the international application This sheet was received by the International Bureau on:
Authorized officer Authorized officer

	52/1	.6	No.
Applicant's or agent's tile B	185 CIP	PCT	International application No.

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

	and to the	murrorganism reiel	rreu to in the description	
on page	28	, line S <u>18</u> -	26	
B. IDENTIFICAT	TION OF DEPOSIT		Further deposits are identified on an additional sheet X	
Name of depositary	nstitution .can Type Cu.	lture Colle	ection	
	v institution (including po			
Rock	Parklawn Di ville, Maryla ed States of	and 20852 America	and Hubmidoma BC G6 AF5	
Identifica	tion Refere	nce by Dep	ositor: Hybridoma, BC.G6.AF5	
Date of deposit 12 Jar	uary 1995 (12.01.95)	HB 11794	
C. ADDITIONA	L INDICATIONS (lea	ve alank if not applicabl	This information is continued on an additional about	
Applicant(s) hereby give notice of my/our intention that samples of the above-identified culture shall be available only to experts in accordance with paragraph 3 of the Fourth Schedule to the Patents Rules 1995.				
D. DESIGNATE	D STATES FOR WE	IICH INDICATIO	ONS ARE MADE (if the indications are not for all designated States)	
Singa				
E. SEPARATE	FURNISHING OF I	NDICATIONS (los	ve blank if not applicable)	
The indications lis Number of Deposit')	ied below will be submitt	ed to the Internations	il Bureau later (specify the general nature of the indications a.g., "Accession	
F	or receiving Office use of	nly -	For international Buresu use only	
	as received with the inte		This sheet was received by the international Bureau on:	
Authorized office	•		Authorized afficer	
1				

52/17

Applicant's or agent's tile reference number	B185	CIP	PCT	International application No.

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made nellow relate to the microorganism referred to in the description on page					
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet 🗵				
Name of depositary institution American Type Culture Collection					
Address at depositary institution traciming postal code and country					
12301 Parklawn Drive Rockville, Maryland 20852 United States of America					
Identification Reference by De					
Date of deposit 12 January 1995 (12.01.95)	Accession Number HB 11795				
C. ADDITIONAL INDICATIONS (leave blank if not applicable	(c) This information is continued on an additional about				
Applicant(s) hereby give notice of my/our intention that samples of the above-identified culture shall be available only to experts in accordance with paragraph 3 of the Fourth Schedule to the Patents Rules 1995.					
D. DESIGNATED STATES FOR WHICH INDICATION	INS ARE MADE (if the indications are not for all danignated States)				
Singapore					
E. SEPARATE FURNISHING OF INDICATIONS (learn	e blank if not applicable)				
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications a.g., "Accusion Number of Deposit")					
For receiving Office use only	For International Bureau use only				
This sheet was received with the international application	This sheet was received by the international Bureau on:				
Authorized officer	Authorized officer				

52/18

Applicant's or agent's file				International application No.
Applicant's or agent's file reference number	B185	CIP	PCT	

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made nellow relate to the microorganism referred to in the description on page 28 time S_18-26				
B. IDENTIFICATION OF DEPOSIT Further deposits are identified on an additional abeet X				
Name of depositary institution American Type Culture Collection				
Address of depositary institution functioning postal code and country)				
12301 Parklawn Drive Rockville, Maryland 20852 United States of America				
Identification Reference by Depositor: Hybridoma, AG.H1.5.1 Date of deposit				
12 January 1995 (12.01.95) Accession Number HB 11796				
C. ADDITIONAL INDICATIONS (Leave niank if not applicable) This information is continued on an additional about				
Applicant(s) hereby give notice of my/our intention that samples of the above-identified culture shall be available only to experts in accordance with paragraph 3 of the Fourth Schedule to the Patents Rules 1995.				
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)				
Singapore				
E. SEPARATE FURNISHING OF INDICATIONS (loave blank if not applicable)				
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Accessed Number of Depart")				
For International Bureau use only				
This sheet was received with the international application This sheet was received by the International Bureau on:				
Authorized officer Authorized officer				

52/19

Applicant's or agent's file reference number	B185	CIP	PCT	International application No.	

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

the same of the sa				
A. The indications made nellow relate to the microorganism referred to in the description on page				
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet			
Name of depositary institution American Type Culture Col	llection			
Address of depositary institution finctuaine postal code and count	(vi)			
12301 Parklawn Drive Rockville, Maryland 20852 United States of America				
Identification Reference by D				
Date of deposit 12 January 1995 (12.01.95	Accession Number HB 11797			
C. ADDITIONAL INDICATIONS (leave plank if not applied	shie) This information is continued on an additional sheet			
Applicant(s) hereby give notice samples of the above-identified only to experts in accordance Fourth Schedule to the Patents	d culture shall be available			
D. DESIGNATED STATES FOR WHICH INDICATI	ONS ARE MADE (if the indications are not for all designated States)			
Singapore				
E. SEPARATE FURNISHING OF INDICATIONS (Les	ve blank if not applicable)			
The indications listed below will be submitted to the internations Number of Deposit*)	l Bureau later (specify the general nature of the indications a.g., "Accommon			
For receiving Office use only	For International Bureau use only			
This sheet was received with the international application	This sheet was received by the International Bureau on:			
Authorized officer	Authorized officer			

_	2	•	2	Λ
כ	L	/	L	u

	52/20	
Applicant's or agent's file reference number	B185 CIP PCT	International application No.

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

	THE UP THE RESCRIPTION
The indications made nellow relate to the microorganism reter on page	
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet X
Name of depositary institution American Type Culture Colle	ection
Address of acpositary institution (including poster code and country)	
12301 Parklawn Drive Rockville, Maryland 20852 United States of America Identification Reference by Dep	oositor: Hybridoma, BD.A8.AB9 Accession Number HB 11798
12 January 1995 (12.01.95)	
C. ADDITIONAL INDICATIONS (Icove plant if not applicable	This information is continued on an additional sheet
Applicant(s) hereby give notice samples of the above-identified only to experts in accordance w Fourth Schedule to the Patents	culture shall be available ith paragraph 3 of the Rules 1995.
D. DESIGNATED STATES FOR WHICH INDICATIO	NS ARE MADE (if the indications are not for all designated States)
Singapore	
E. SEPARATE FURNISHING OF INDICATIONS (loss)	e blenk if not applicable)
The indications listed below will be submitted to the International Number of Deposit*)	Bureau later (specify the general nature of the indicamental a.g., Accessed
This sheet was received with the international application	
Authorized officer	Authorized efficer

52/21

	JE/ ZI	
Applicant's or agent's file		International application No.
reterence number	B185 CIP PCT	

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

A. The indications made below relate to the microorganism retion page 28 , line S 18	erreu to in the description - 26
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional about
Name of depositary institution American Type Culture Coll	ection
Address of depositary institution tracing posts code and country	
12301 Parklawn Drive Rockville, Maryland 20852 United States of America	
	positor: Hybridoma, BK.All.AC10
Date of deposit 12 January 1995 (12.01.95)	HB 11799
C. ADDITIONAL INDICATIONS (Icave signit if not applicable	(c) This information is continued on an additional sheet
Applicant(s) hereby give notice samples of the above-identified only to experts in accordance we Fourth Schedule to the Patents	d culture shall be available
D. DESIGNATED STATES FOR WHICH INDICATION	INS ARE MADE (if the indications are not for all designated States)
Singapore	
E. SEPARATE FURNISHING OF INDICATIONS (loss)	e blank if not applicable)
The indications listed below will be submitted to the International Number of Deposit*)	Bureau later (specify the general noture of the indications a.g., "Accesses
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	This sheet was received by the International Bureau on:
Authorized officer	Authorized officer

PCT/US96/01386

What is claimed is:

WO 96/22788

- 1. A method for treating or reducing the advancement, severity or effects of neoplasia comprising the step of administering a therapeutically effective amount of a LT- α/β heteromeric complex and a pharmaceutically acceptable carrier.
 - 2. The method according to claim 1, wherein the LT- α/β heteromeric complex has a LT- $\alpha1/\beta2$ stoichiometry.
- The method according to claim 1, wherein the LT-α/β
 heteromeric complex is a soluble LT-α/β heteromeric complex.
- The method according to any one of claims 1-3, wherein the LT-α subunit is selected from the group consisting of lymphotoxin-α, native human or animal lymphotoxin-α, recombinant lymphotoxin-α, soluble lymphotoxin-α, secreted lymphotoxin-α, lymphotoxin-α muteins, or lymphotoxin-α-active fragments of any of the above.
- 5. The method according to any one of claims 1-3,
 wherein the LT-ß subunit is selected from the group
 consisting of lymphotoxin-ß, native human or animal
 lymphotoxin-ß, recombinant lymphotoxin-ß, soluble
 lymphotoxin-ß, secreted lymphotoxin-ß, lymphotoxin-ß
 muteins, or lymphotoxin-ß-active fragments of any of the
 above.
 - 6. The method according to claim 3, wherein the LT-ß subunit is cleaved between amino acids 44 and 88 and the N-terminal portion replaced with a type I leader sequence.

- 54 -

- 7. The method according to claim 6, wherein the type I leader sequence is the vascular cell adhesion molecule 1 (VCAM-1) leader sequence.
- The method according to any one of claims 1-3,
 wherein the LT-α/β heteromeric complex is administered in the presence of a therapeutically effective amount of at least one LT-β-R activating agent.
- 9. The method according to claim 8, wherein one LT-B-R

 10 activating agent comprises a therapeutically effective

 amount of IFN-y.
 - 10. The method according to claim 9, wherein a second LT-B-R activating agent comprises a therapeutically effective amount of an anti-LT-B-R antibody.
- 15 11. The method according to claim 10, wherein the anti-LT-B-R antibody is a monoclonal antibody.

ť,

20

- 12. The method according to claim 11, wherein the anti-LT-B-R monoclonal antibody is selected from the group consisting of anti-LT-B-R mAb BKA11, CDH10, BCG6 and BHA10.
 - 13. A method for treating or reducing the advancement, severity or effects of neoplasia comprising the step of administering a therapeutically effective amount of at least two LT-B-R activating agents and a pharmaceutically acceptable carrier.
 - 14. The method according to claim 13, wherein at least one LT- β -R activating agent comprises an anti-LT- β -R antibody.

- 55 -

- 15. The method according to claim 14, wherein the anti-LT- β -R antibody is CBE11.
- 16. The method according to claim 13, wherein the LT-β-R activating agents comprise at least two anti-LT-β-R monoclonal antibodies which are directed against non-overlapping epitopes of LT-β-R.
- 17. The method according to claim 16, wherein one anti-LT-ß-R monoclonal antibody is selected from the group consisting of AGH1 and BDA8, and another anti-LT-ß-R monoclonal antibody is selected from the group consisting of BCG6, BHA10, BKA11, CDH10 and CBE11.
- 18. The method according to claim 16, wherein one anti-LT-ß-R monoclonal antibody is selected from the group consisting of BCG6 and BHA10, and another anti-LT-ß-R monoclonal antibody is selected from the group consisting of AGH1, BDA8, BKA11, CDH10 and CBE11.
- 19. The method according to claim 16, wherein one anti-LT-\beta-R monoclonal antibody is selected from the group consisting of BKA11 and CDH10, and another anti-LT-\beta-R 20 monoclonal antibody is selected from the group consisting of AGH1 and BDA8, BCG6, BHA10, and CBE11.
 - 20. The method according to claim 16, wherein one anti-LT-B-R monoclonal antibody is CBE11, and another anti-LT-B-R monoclonal antibody is selected from the group consisting of AGH1, BDA8, BCG6, BHA10, BKA11, CDH10 and CBE11.

25

21. The method according to claim 16, wherein the anti-LT- β -R monoclonal antibodies are CBE11 and BHA10.

- 56 -

- 22. The method according to claim 16, wherein the anti-LT-B-R monoclonal antibodies are CBE11 and CDH10.
- 23. The method according to claim 16, wherein the anti-LT-ß-R monoclonal antibodies are AGH1 and CDH10.
- 5 24. The method according to any one of claims 13-23, wherein one LT- β -R activating agent is IFN- γ .
 - 25. A method for treating or reducing the advancement, severity or effects of neoplasia comprising the step of administering a therapeutically effective amount of cross-linked anti-LT-B-R antibodies as a first LT-B-R
- cross-linked anti-LT-ß-R antibodies as a first LT-ß-activating agent in the presence of a second LT-ß-R activating agent and a pharmaceutically acceptable carrier.
- 26. The method according to claim 25, wherein the crosslinked anti-LT-B-R antibodies are non-covalently immobilized on a surface.
 - 27. The method according to claim 25, wherein the cross-linked anti-LT-B-R antibodies are covalently immobilized on a surface.
- 28. The method according to claim 25, wherein the cross-linked anti-LT-B-R antibodies are non-covalently aggregated in solution by means of an anti-LT-B-R antibody cross-linking agent.
- 29. The method according to claim 28, wherein the anti-25 LT-B-R antibody cross-linking agent comprises a secondary antibody directed against the anti-LT-B-R antibody.

- 30. The method according to claim 28, wherein the anti-LT-B-R antibody cross-linking agent comprises an Fc receptor which binds to the anti-LT-B-R antibody.
- 31. The method according to claim 25, wherein the cross-linked anti-LT-B-R antibodies are covalently aggregated in solution by means of a chemical anti-LT-B-R antibody cross-linking agent.
- 32. The method according to any one of claims 25-31, wherein the second LT-B-R activating agent comprises 10 IFN-y.

- 33. A method for treating or reducing the advancement, severity or effects of neoplasia comprising the step of administering a therapeutically effective amount of at least one LT-B-R activating agent and a pharmaceutically acceptable carrier.
 - 34. The method according to claim 33, wherein at least one LT- β -R activating agent comprises an anti-LT- β -R antibody.
- 35. The method according to claim 34, wherein the anti-20 LT-B-R antibody is CBEll.
 - 36. A method for selecting a LT-B-R activating agent which acts in the presence of LT- α/B heteromeric complexes comprising the steps of:
- a) culturing tumor cells in the presence of
 25 LT-α/β heteromeric complexes, an effective amount of a first LT-β-R activating agent and a second putative
 LT-β-R activating agent; and
 - b) determining whether the second putative LT-B-R activating agent increases the anti-tumor activity

of the LT- α/β heteromeric complex in the presence of the first LT- β -R activating agent.

- 58 -

- 37. The method according to claim 36, wherein the first LT- β -R activating agent is IFN- γ .
- 5 38. The method according to claim 36, wherein the LT- α/β heteromeric complex has a LT- α/β 2 stoichiometry.
 - 39. A pharmaceutical composition comprising a therapeutically effective amount of a LT- α/β heteromeric complex and a pharmaceutically acceptable carrier.
- 10 40. The pharmaceutical composition according to claim 39, wherein the LT- α/β heteromeric complex has a LT- $\alpha1/\beta2$ stoichiometry.
 - 41. The pharmaceutical composition according to claim 39, wherein the LT- α/β heteromeric complex is soluble.
- 15 42. The pharmaceutical composition according to any one of claims 39-41, further comprising a therapeutically effective amount of at least one LT-B-R activating agent.
 - 43. The pharmaceutical composition according to claim
 - 42, wherein one LT-B-R activating agent is IFN-y.
- 44. The pharmaceutical composition according to claim42, wherein one LT-β-R activating agent is an anti-LT-β-R
 - 45. The pharmaceutical composition according to claim
 - 44, wherein the anti-LT-B-R antibody is a monoclonal
- 25 antibody.

antibody.

- 46. The pharmaceutical composition according to claim 45, wherein the anti-LT-B-R monoclonal antibody is selected from the group consisting of anti-LT-B-R mAb BKA11, CDH10, BCG6, and BHA10.
- 5 47. A pharmaceutical composition comprising a therapeutically effective amount of at least two LT- β -R activating agents without exogenous LT- α/β heteromeric complex, and a pharmaceutically acceptable carrier.
- 48. The pharmaceutical composition according to claim
 10 47, wherein at least one LT-B-R activating agent
 comprises an anti-LT-B-R antibody.
 - 49. The pharmaceutical composition according to claim 48, wherein the anti-LT-B-R antibody is a monoclonal antibody.
- 15 50. The pharmaceutical composition according to claim 49, wherein the anti-LT-B-R monoclonal antibody is CBE11.
 - 51. The pharmaceutical composition according to claim 47, wherein at least two LT-ß-R activating agents comprise anti-LT-ß-R monoclonal antibodies which are directed against non-overlapping epitopes of LT-ß-R.

20

52. The pharmaceutical composition according to claim 51, wherein one anti-LT-\$BR monoclonal antibody is selected from the group consisting of AGH1 and BDA8, and another anti-LT-\$BR monoclonal antibody is selected from the group consisting of BCG6, BHA10, BKA11, CDH10 and CBE11.

5

- 53. The pharmaceutical composition according to claim 51, wherein one anti-LT-ß-R monoclonal antibody is selected from the group consisting of BCG6 and BHA10, and another anti-LT-ß-R monoclonal antibody is selected from the group consisting of AGH1, BDA8, BKA11, CDH10 and CBE11.
- 54. The pharmaceutical composition according to claim 51, wherein one anti-LT-B-R monoclonal antibody is selected from the group consisting of BKA11 and CDH10, and another anti-LT-B-R monoclonal antibody is selected from the group consisting of AGH1 and BDA8, BCG6, BHA10, and CBE11.
- 55. The pharmaceutical composition according to claim 51, wherein one anti-LT-\(\beta\)-R monoclonal antibody is CBE11, and another anti-LT-\(\beta\)-R monoclonal antibody is selected from the group consisting of AGH1, BDA8, BCG6, BHA10, BKA11, CDH10 and CBE11.
 - 56. The pharmaceutical composition according to claim 51, wherein the anti-LT-ß-R monoclonal antibodies are CBE11 and BHA10.
 - 57. The pharmaceutical composition according to claim 51, wherein the anti-LT-ß-R monoclonal antibodies are CBE11 and CDH10.
- 58. The pharmaceutical composition according to claim 25 51, wherein the anti-LT-ß-R monoclonal antibodies are AGH1 and CDH10.
 - 59. The pharmaceutical composition according to any one of claims 51-58, further comprising IFN- γ as one of the LT- β -R activating agents.

- 61 -

- 60. A pharmaceutical composition comprising a therapeutically effective amount of cross-linked anti-LT-B-R antibodies as a LT-B-R activating agent and a pharmaceutically acceptable carrier.
- 5 61. The pharmaceutical composition according to claim 60, wherein the cross-linked anti-LT-B-R antibodies are non-covalently immobilized on a surface.
- 62. The pharmaceutical composition according to claim 60, wherein the cross-linked anti-LT-B-R antibodies are covalently immobilized on a surface.
 - 63. The pharmaceutical composition according to claim 60, wherein the cross-linked anti-LT-\beta-R antibodies are non-covalently aggregated in solution by means of an anti-LT-\beta-R antibody cross-linking agent.
- 15 64. The pharmaceutical composition according to claim 63, wherein the anti-LT-B-R antibody cross-linking agent comprises a secondary antibody directed against the anti-LT-B-R antibody.
- 65. The pharmaceutical composition according to claim
 20 60, wherein the cross-linked anti-LT-B-R antibodies are
 covalently aggregated in solution by means of a chemical
 anti-LT-B-R antibody cross-linking agent.
- 66. The pharmaceutical composition according to any one of claims 60-65, further comprising IFN-γ as a second
 LT-β-R activating agent.
 - 67. A pharmaceutical composition comprising a therapeutically effective amount of at least one LT-B-R

activating agent without exogenous LT- α/β heteromeric complex, and a pharmaceutically acceptable carrier.

- 68. The pharmaceutical composition according to claim 67, wherein at least one LT-ß-R activating agent comprises an anti-LT-ß-R antibody.
 - 69. The pharmaceutical composition according to claim 68, wherein the anti-LT-\$-R antibody is CBE11.
 - 70. An LT- β -R activating agent selected according to the method of claim 36.

Fig. 1A

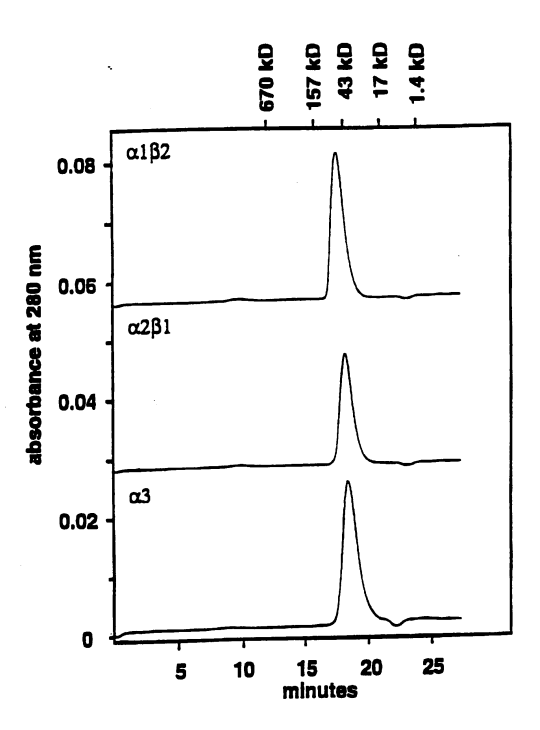
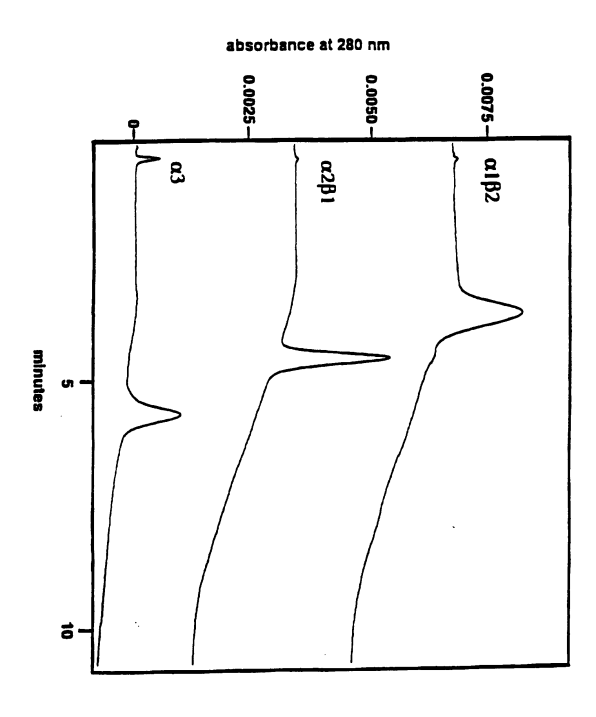


Fig. 1B



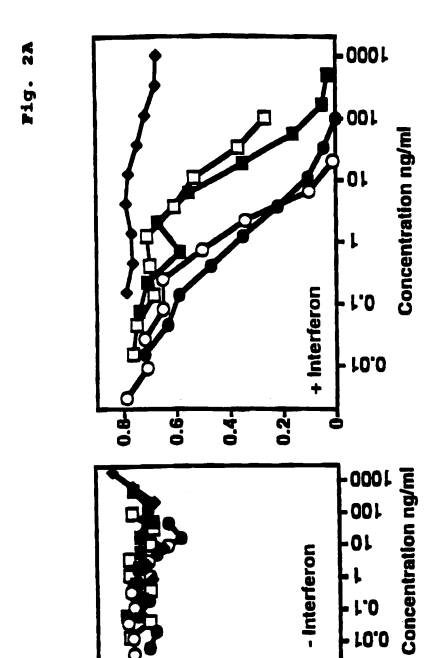
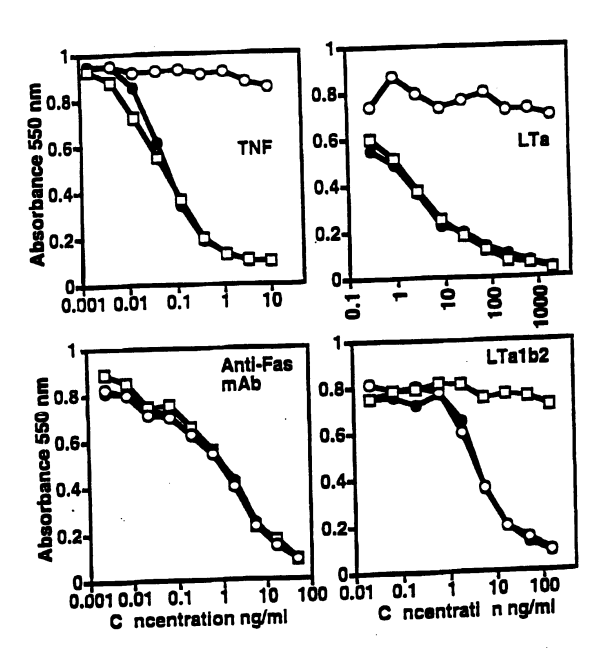
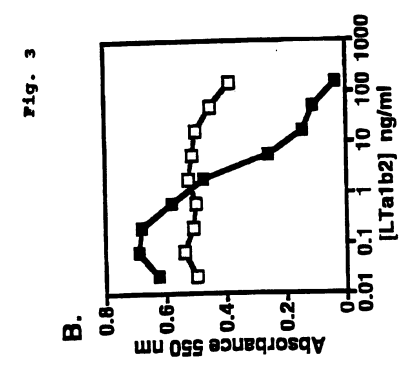
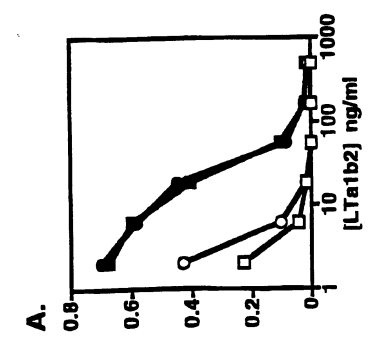
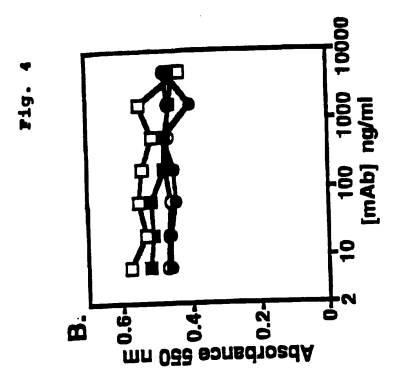


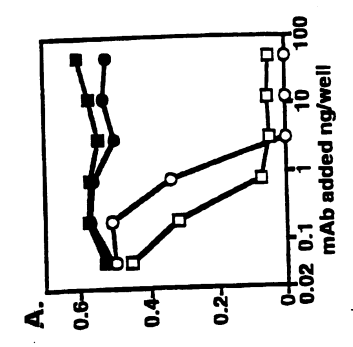
Fig. 2B

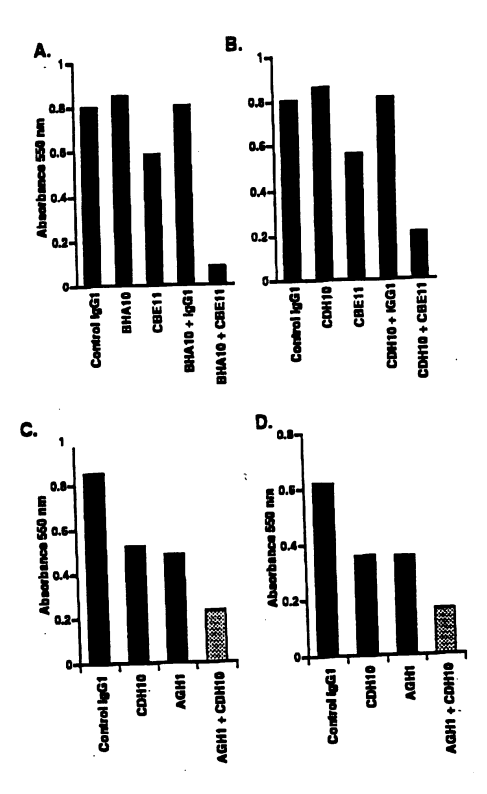


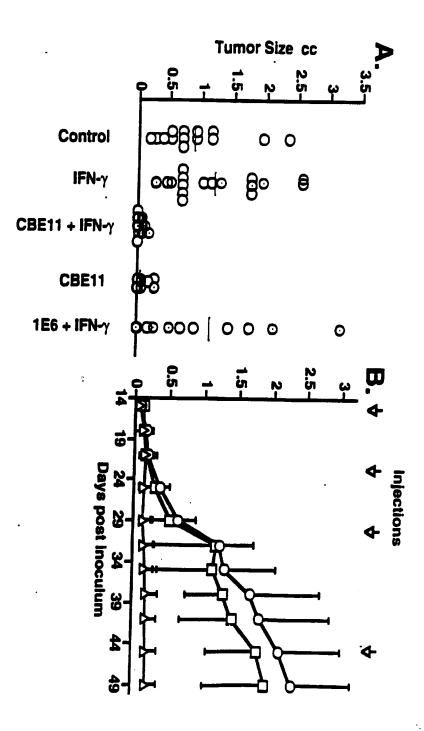












Intr ional Application No PCT/US 96/91386

		PC T/U	S 96/01386
A. CLASSI IPC 6	FICATION OF SUBJECT MATTER A61K38/19 A61K39/395 A61K38/ //(A61K39/395,38:21),(A61K38/21,3	21 G01N33/53 8:19)	
According to	o International Patent Classification (IPC) or to both national class	ification and IPC	
	SEARCHED		
Minimum d IPC 6	ocumentation searched (classification system followed by classification sy	uion symbols)	
Documentat	ion searched other than minimum documentation to the extent that	such documents are included in the	fields searched
Electronic d	ata base consulted during the international search (name of data be	use and, where practical, search term	ns used)
C. DOCUM	IENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
Х	WO,A,94 13808 (BIOGEN INC. ET AL	.) 23 June	1-7, 39-41
	cited in the application see page 33, line 6 - line 12 see page 34, line 31 - page 36,	line 8	
X	WO,A,92 00329 (BIOGEN INC.) 9 Ja cited in the application see claims	nuary 1992	1-7, 39-41
		-/	
	`		
X Furt	her documents are listed in the continuation of box C.	X Patent family members at	re listed in annex.
'A' docum	tegories of cited documents : ent defining the general state of the art which is not cred to be of pericular relevance	"T later document published after or priority date and not in co-cited to understand the principle invention."	milict with the application out
"L" docume which	ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another	"X" document of particular releva cannot be considered novel o involve an inventive step who "Y" document of particular releva	r cannot be considered to in the document is taken alone inor; the claimed invention
'O' docum	n or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but han the priority date claimed	document is combined with o	ive an inventive step when the one or more other such docu- ng obvious to a person skilled ne patent family
Date of the	actual completion of the international search	Date of mailing of the interna 0 7. 06, 96	
	3 May 1996	Authorized officer	
NAMES AND E	European Patent ffice, P.B. 5818 Patentiaan 2 NL - 2280 HV Ripswik Td. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Nooij, F	

Intr ional Application No PCI/US 96/01386

DOCUMENTS CONTINUED TO BE BEI FVANT	PC1/02 30/01300
Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
SCIENCE, vol. 264, no. 5159, 29 April 1994, WASHINGTON, DC, USA, pages 707-710, XP002003806 P. CROWE ET AL.: "A lymphotoxin-beta-specific receptor." cited in the application see page 708, left-hand column, line 40 - line 62 see page 708, middle column, line 31 - might-hand column, line 28	39-41
right-hand corumns, The 20	1.70
CELL, vol. 72, 26 March 1993, CAMBRIDGE, MA, USA, pages 847-856, XP002003807 J. BROWNING ET AL.: "Lymphotoxin beta, a novel member of the TNF family that forms a heteromeric complex with lymphotoxin on the cell surface." cited in the application see abstract	1-70
THE JOURNAL OF IMMUNOLOGY, vol. 154, no. 1, 1 January 1995, BALTIMORE, MD, USA, pages 33-46, XP002003808 J. BROWNING ET AL.: "Characterization of surface lymphotoxin forms. Use of specific monoclonal antibodies and soluble receptors." cited in the application see abstract see discussion	1-70
US,A,4 770 995 (RUBIN ET AL.) 13 September 1988 see examples 2,6,7 see claims	36-38
THE FASEB JOURNAL, vol. 8, no. 5, 18 March 1994, BETHESDA, MD, USA, page A962 XP002003809 C. WARE ET AL.: "A novel lymphotoxin beta (LTbeta) specific receptor and TNF receptors identify distinct surface LTalpha/beta complexes." see abstract 5571	1-70
	SCIENCE, vol. 264, no. 5159, 29 April 1994, WASHINGTON, DC, USA, pages 707-710, XP002003806 P. CROWE ET AL.: "A lymphotoxin-beta-specific receptor." cited in the application see page 708, left-hand column, line 40 - line 62 see page 708, middle column, line 31 - right-hand column, line 28 CELL, vol. 72, 26 March 1993, CAMBRIDGE, MA, USA, pages 847-856, XP002003807 J. BROWNING ET AL.: "Lymphotoxin beta, a novel member of the TNF family that forms a heteromeric complex with lymphotoxin on the cell surface." cited in the application see abstract THE JOURNAL OF IMMUNOLOGY, vol. 154, no. 1, 1 January 1995, BALTIMORE, MD, USA, pages 33-46, XP002003808 J. BROWNING ET AL.: "Characterization of surface lymphotoxin forms. Use of specific monoclonal antibodies and soluble receptors." cited in the application see abstract see discussion US,A,4 770 995 (RUBIN ET AL.) 13 September 1988 see examples 2,6,7 see claims THE FASEB JOURNAL, vol. 8, no. 5, 18 March 1994, BETHESDA, MD, USA, page A962 XP002003809 C. WARE ET AL.: "A novel lymphotoxin beta (LTbeta) specific receptor and TNF receptors identify distinct surface

Intr onal Application No PCI/US 96/01386

		PC1/US 96/01386
C./Continu	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
P,X	F. MACKAY ET AL.: "Lymphotoxin-beta receptor triggering induces activation of the NF-kB transcription factor in various cell types. In: 'THE 9th INTERNATIONAL CONGRESS OF IMMUNOLOGY, San Francisco, July 23-29, 1995. Abstract book." July 1995, SAN FRANCISCO, CA, USA XP002003810 see abstract 4583	1-8, 13-23, 25-31, 33-35, 39-42, 44-58, 60-65, 67-70
P,X	J. BROWNING ET AL.: "Signalling through the lymphotoxin-beta receptor in conjunction with interferon-gamma induces the death of a human tumor line. In: THE 9th INTERNATIONAL CONGRESS OF IMMUNOLOGY, San Francisco, July 23-29, 1995. Abstract book." July 1995, SAN FRANCISCO, CA, USA XP002003811 see abstract 4582	1-35, 39-70
T	THE JOURNAL OF EXPERIMENTAL MEDICINE, vol. 183, no. 3, 1 March 1996, NEW YORK, NY, USA, pages 867-878, XP000571445 J. BROWNING ET AL.: "Signalling through the lymphotoxin beta receptor induces the death of some adenocarcinoma tumor lines." see the whole document	1-70

Information on patent family members

Int ional Application No PCT/US 96/01386

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
		AU-B- CA-A- EP-A-	6048394 2150249 0672143	04-07-94 23-06-94 20-09-95
WO-A-9200329	09-01-92	AU-B- AU-B- CA-A- EP-A- JP-T-	2716795 8228891 2086264 0536299 6501456	15-02-96 23-01-92 28-12-91 14-04-93 17-02-94
US-A-4770995	13-09-88	NONE		